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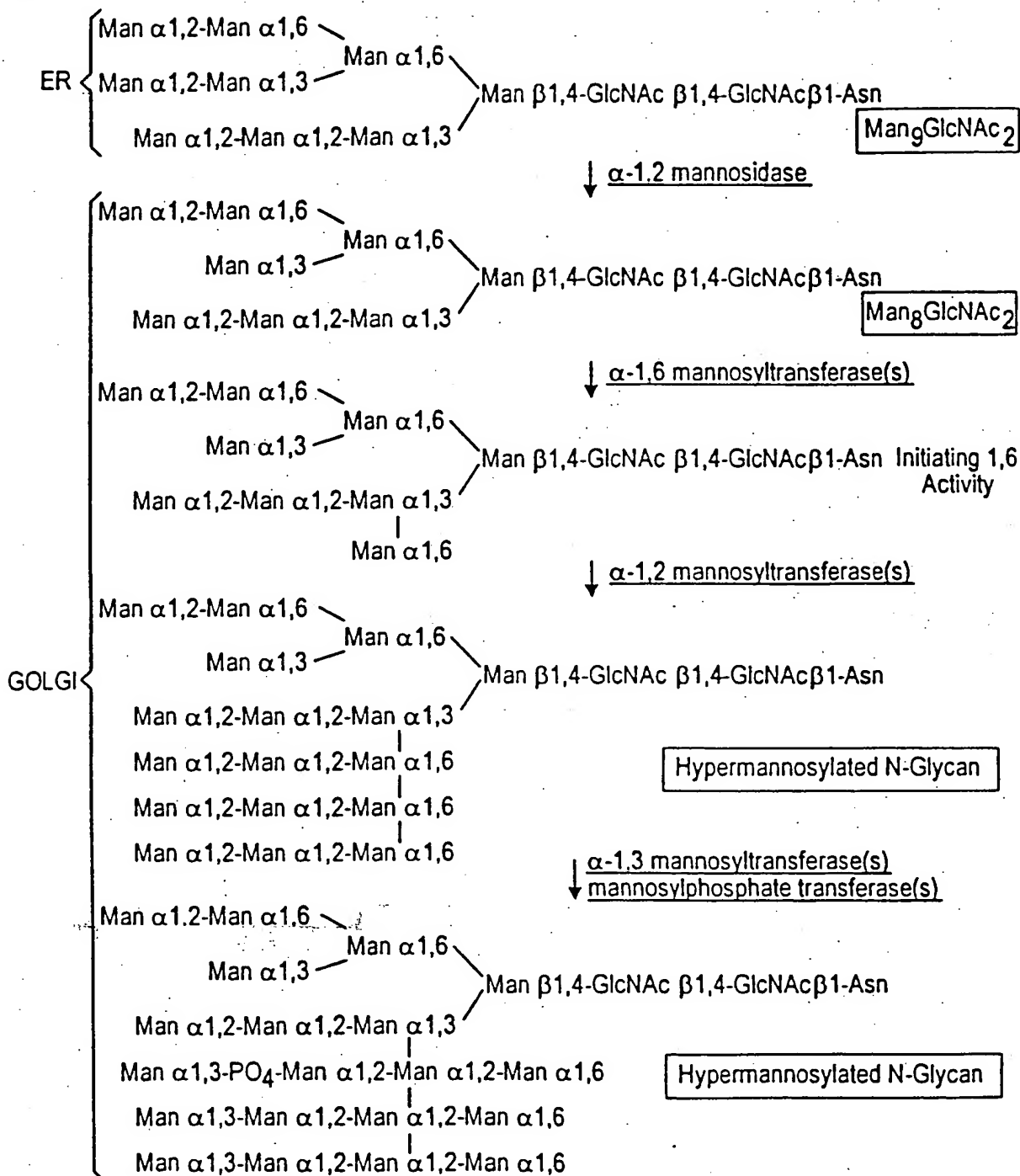


FIG. 1A

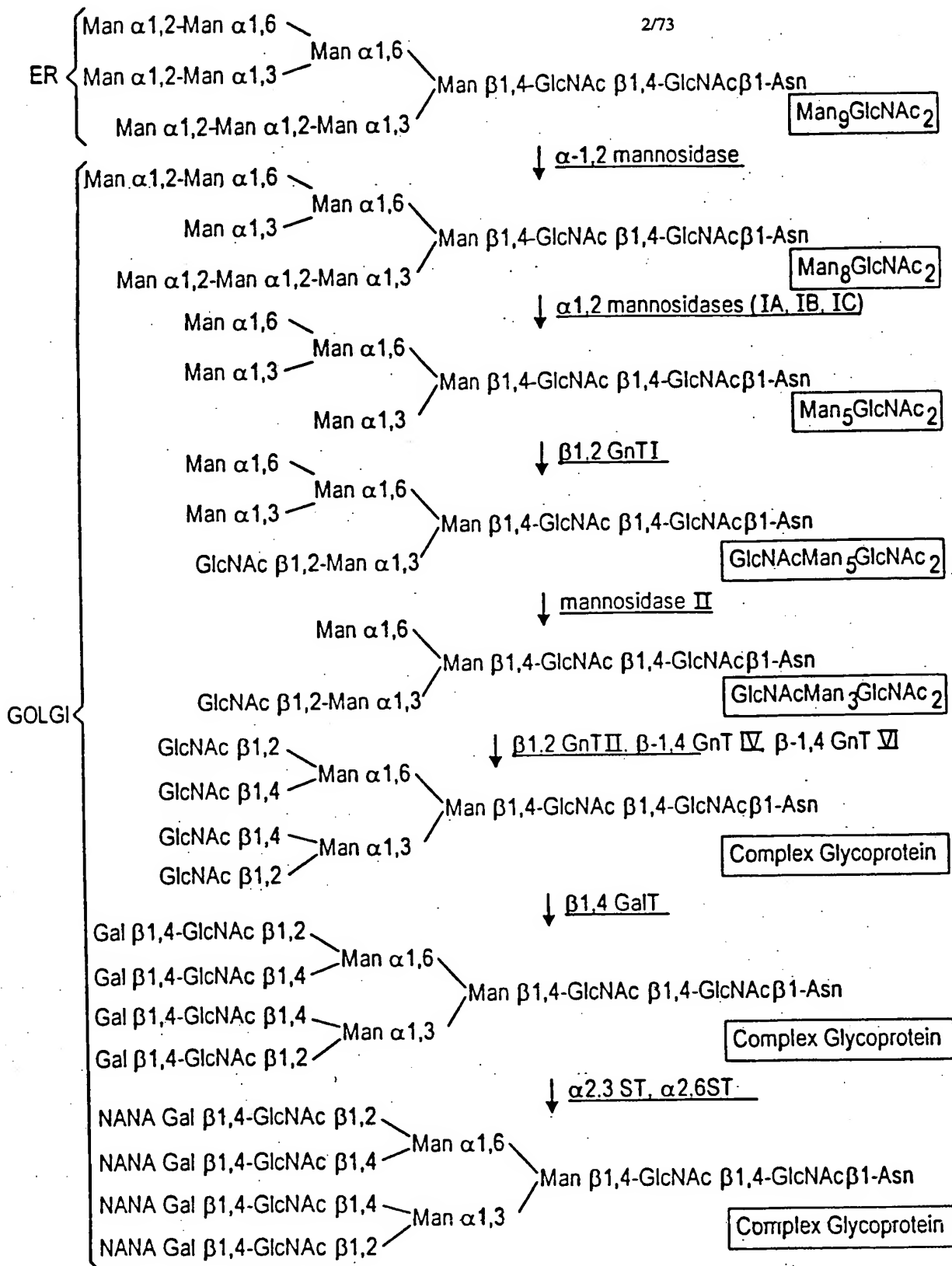


FIG. 1B

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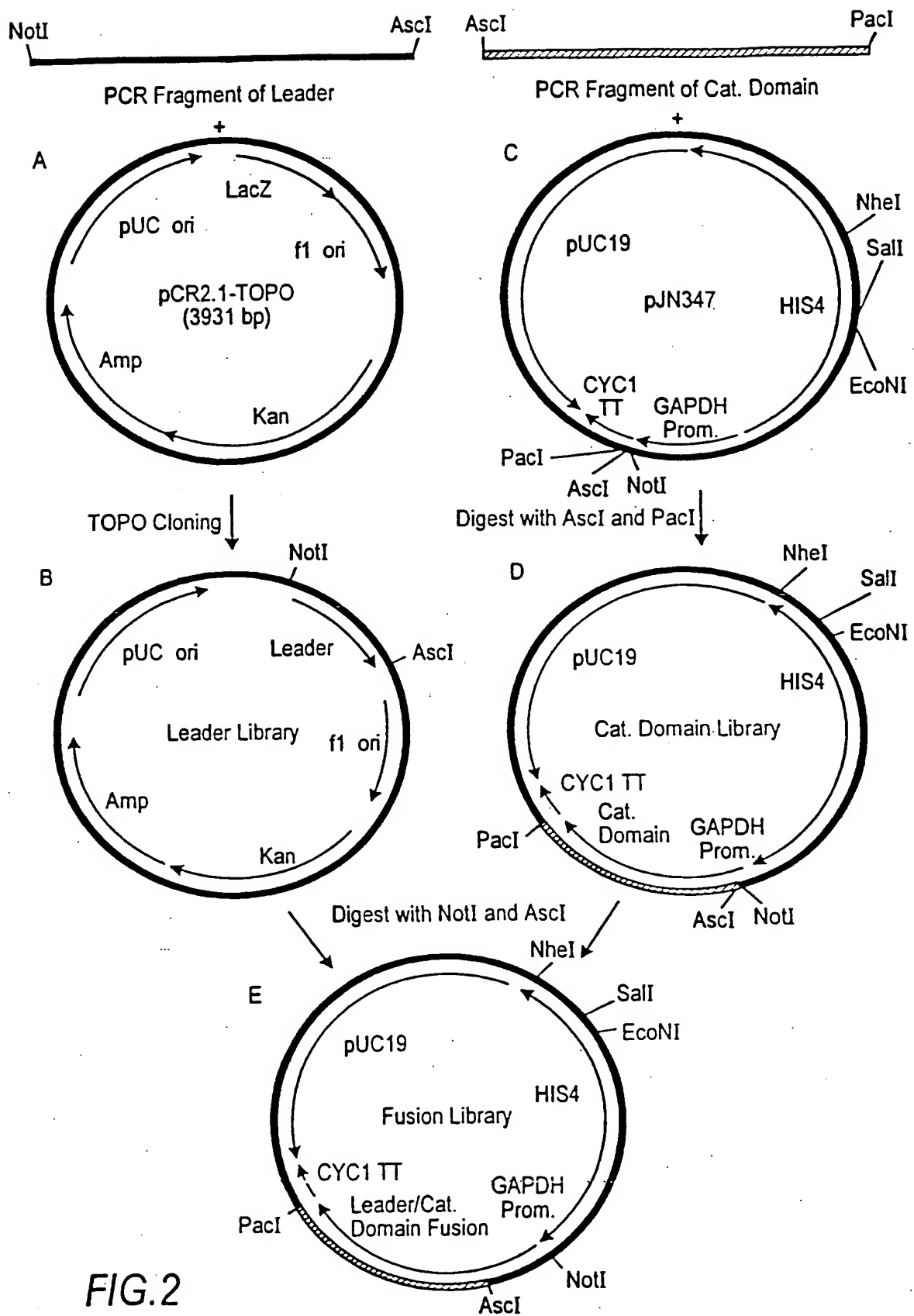


FIG.2

**M. musculus alpha-1,2-mannosidase IA open reading frame.** The transmembrane and catalytic domains are highlighted in bold respectively. The sequence of the primers used to generate the N-terminal truncations are highlighted by underlining and the start of each respective protein fragment indicated by an arrow.

1 atgcccgtaggggacctgttgcgcgtcttcagtaccctggggcgccctgggcagtgccctgggcgggggcttggcggcgaggaggaagggg  
11 M P V G G L L P L F S S P G G G L G S G L G G L G G L G G R K G  
97 tctgccccgcttcgcctcaccgagaagtctgtctgtctgtgttcagcgccttcacgcgtctgtctgttcggggcaatc  
33 S G P A A F R L T E K F V L L L V F S A F I T L C F G A I  
184 ttctctcgtcgtgactcctccaagtctcagcgggtcctgttcactccaacctgccttcagccgcggcgagacaaagcccggtctcg  
62 F F L P D S S K L L S G V L F H S N P A L Q P P A E H K P G L  
d65 primer  
270 ggcgcgtgcggagatccgcgcgaggggagagtcgcgcacccgaggaaggcgccctgggaccctggagctggactggaagacaacttagcca  
93 G A R A E D A A E G R V R H R E E G A P G D P G A G L E D N L A  
d105 primer  
374 ggcacccgcaaaaccacgagcggtctcaggaagcccaaggagaccctgcagaagctccggaggagatccaaagagacattctgctggagaag  
125 R I R E N H E R A L R E A K E T L Q K L P E E I Q R D I L L E K  
470 aaaagtgccccagaccagctgcgtgacaaggatctgttagggcttgcccaagtggaacttctgccccctgcgggttagagaacccgggagc  
157 E K V A Q D Q L R D K D L F R G L P K V D F L P P V G V E N R E  
d187 primer  
566 ccgctgacccaccatccgtggaagagggcaagatcaagagatgatgacccatgcttggaaataattataacgctatgcgctggggg  
189 P A D A T I R E K R A K I K E M M T H A W N N Y K R Y A W G

**FIG. 3**

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655 ttgaacgaactgaacccctatatcaaaagaaggccattcaagcagtttgtttgggaacatcaaaaggagctacaatagtagatg  
219▶ L N E L K P I S K E G H S S S L F G N I K G A T I V D  
737 cctggatacccttttcattatgggcagatgaagactgaatttcaagaagctaaatcggtgattaaanaatttagatttttaa  
246▶ A L D T L F I M G M K T E F Q E A K S W I K K Y L D F N  
819 tgtgaatgctgaagtcttctgttttgaagtaacatacagcttcgtcggtggaotgtgcagcctactatttgcgcggagag  
273▶ V N A E V S V F E V N I R F V G G L L S A Y Y L S G E  
901 gagatatctcgaagaagcagtggaacttgggttaaaattgctacotgcatttcaactcctcctcctggaataccttgggcac  
301▶ E I F R K K A V E L G V K L L P A F H T P S G I P W A  
983 tgcetgaatatgaagaagtggatcgggcggaactggccctggccctctggaggcagcagatcctcctggccgaatttggaaactct  
328▶ L L N M K S G I G R N W P W A S G G S S I L A E F G T L  
1065 gcatttagagtttatgcacttgcctccttatcaggagaccagctcttgcggaaaagggttatgaaaattcgaacagtggtg  
355▶ H L E F M H L S H L S G D P V F A E K V M K I R T V L  
1147 acaaaactggacaaaaccagaaggccctttatcctaaactatctgaacccacgtgtagcagtggtgggtcacaatcctgtgtcgg  
383▶ N K L D K P E G L Y P N Y L N P S S G Q W G Q H H V S  
1229 ttgaggacttgagacagcttttatgaatatattgtttaaggcgtggttaattgtctgacaagacagatctcgaagcgaagaa  
410▶ V G G L G D S F Y E Y L L K A W L M S D K T D L E A K K  
1311 gatgtatttgatgtgttcaggccatcgagactcactgtgatcgcgaagtcaagtggtgggactaaactacgtacatcgacagtggtg  
437▶ M Y F D A V Q A I E T H L I R K S S G G L T Y I A E W  
1393 aagggggcccctcctggaaacacacagatgggcccacctgacgtgcttgcaggaggcactgtttgcacttggggcagatggagctc  
465▶ K G G L L E H K M G H L T C F A G G M F A L G A D G A  
1475 cggaagcccgcccaacactaccttgaaactcggagctgaatggccgcactgtctcatgatttataatcgtacatatgt  
492▶ P E A R A Q H Y L E L G A E I A R T C H E S Y N R T Y V  
1557 gaagtgggaccggaagcgtttcgatttgatggcgtgtggaagctattggccagggcgaataatgaaaagtattacatttta  
519▶ K L G P E A F R F D G G V E A I A T R Q N E K Y Y I L  
1639 cggcccgggtcatcgagacatacatgtacatgttgccgactgaactcagaccccaagtaacaggacactggccctgggaagccg  
547▶ R P E V I E T Y M Y M W R L T H D P K Y R T W A W E A  
1721 tggaggctctagaagtcactcagagatgaacggaggctactcaggcttaagggtgtttacattggccggtgagagtatga  
574▶ V E A L E S H C R V N G G Y S G L R D V Y I A R E S Y D  
1803 cgatgtccagcaaaagtcttctcctggcagagacactgaagtatttgcacttgatatttccgatgatgacotttctccatba  
601▶ D V Q Q S F F L A E T L K Y L Y L I F S D D L L P L  
1885 gaacactggatcttcaacacccgaggtcctcctcctatactcctgtaacagaaggaattggtggcaagagaatga  
629▶ E H W I F N T E A H P F P I L R E Q K K E I D G K E K

FIG. 3 CONT

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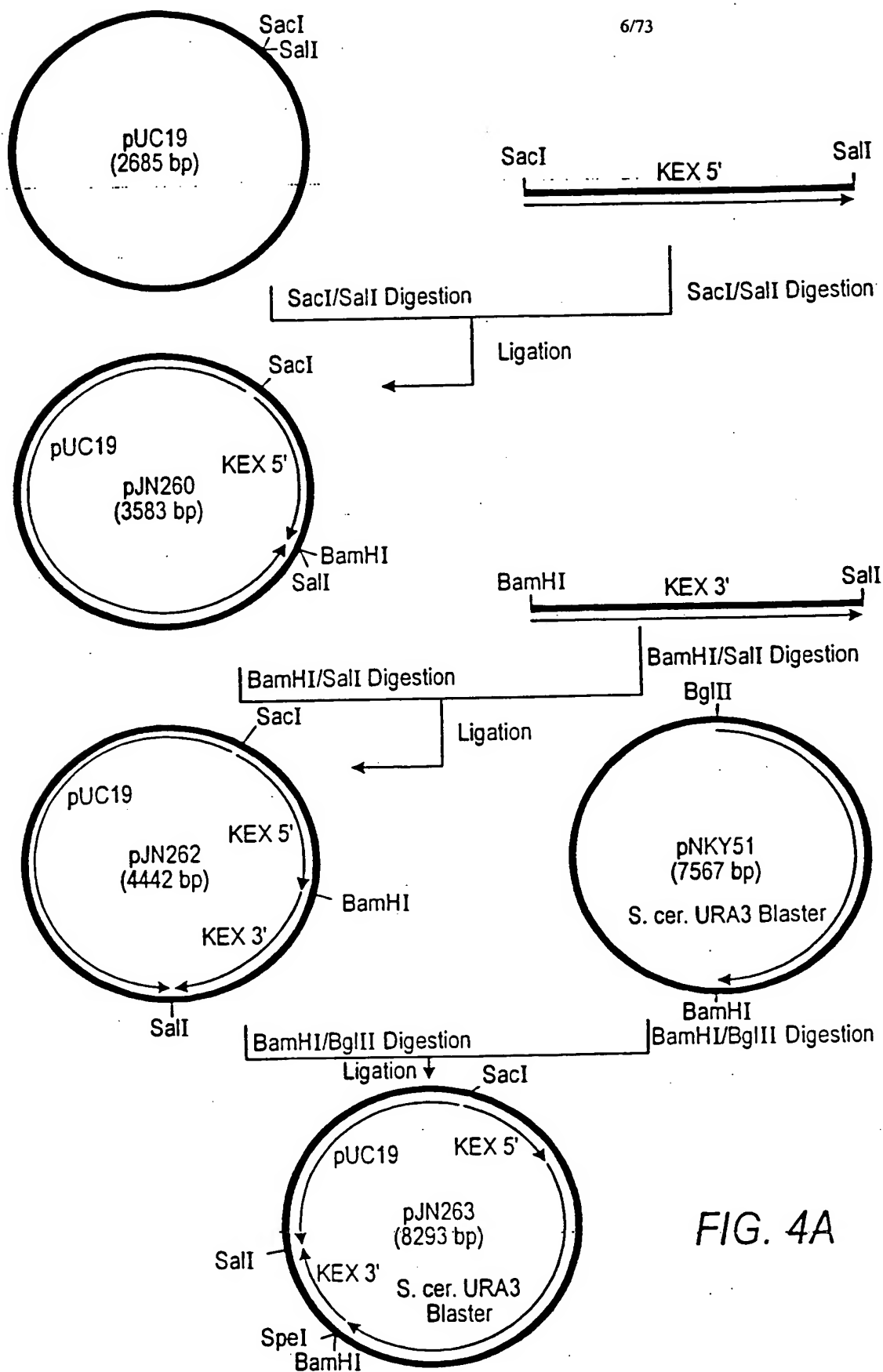


FIG. 4A

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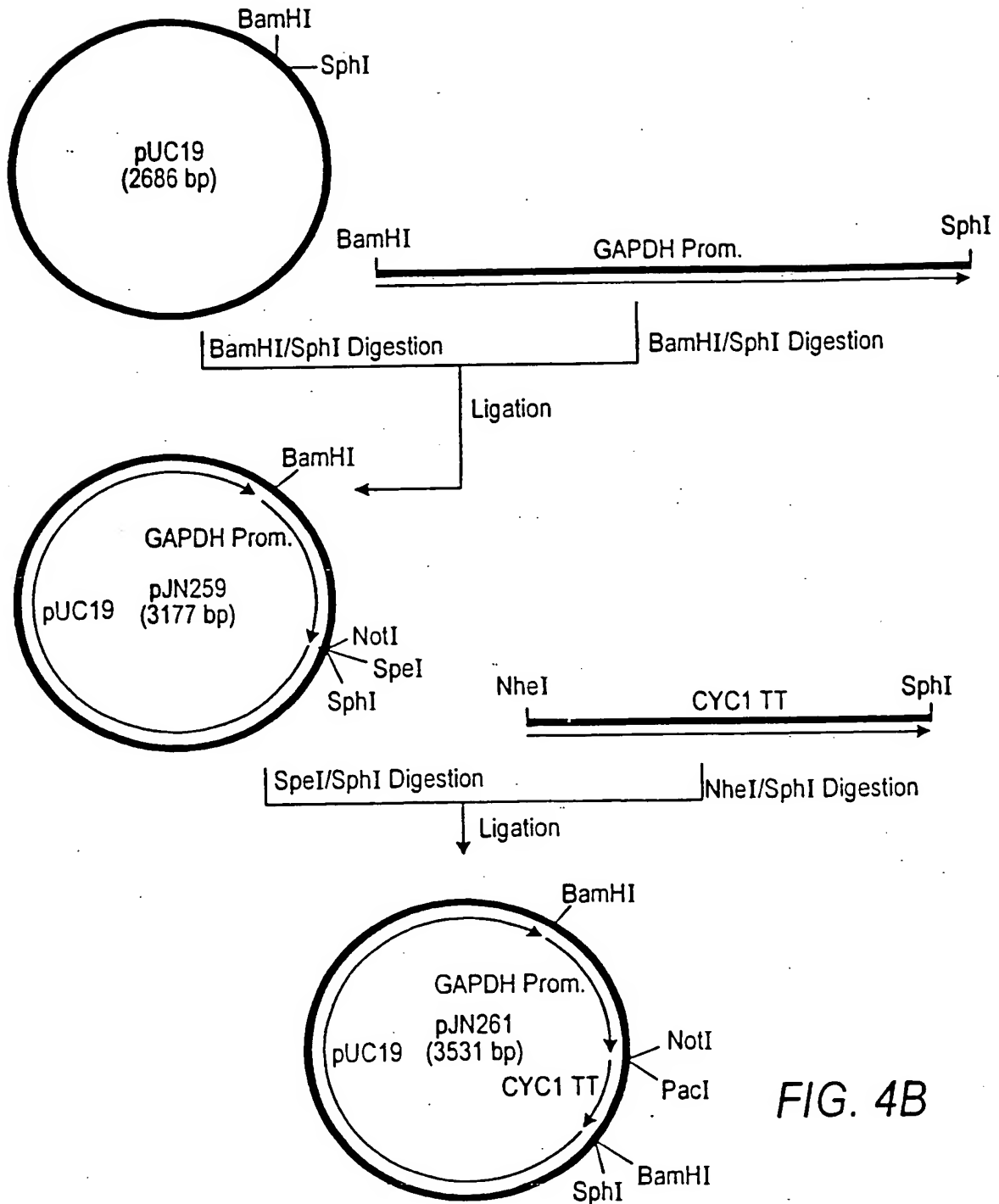


FIG. 4B

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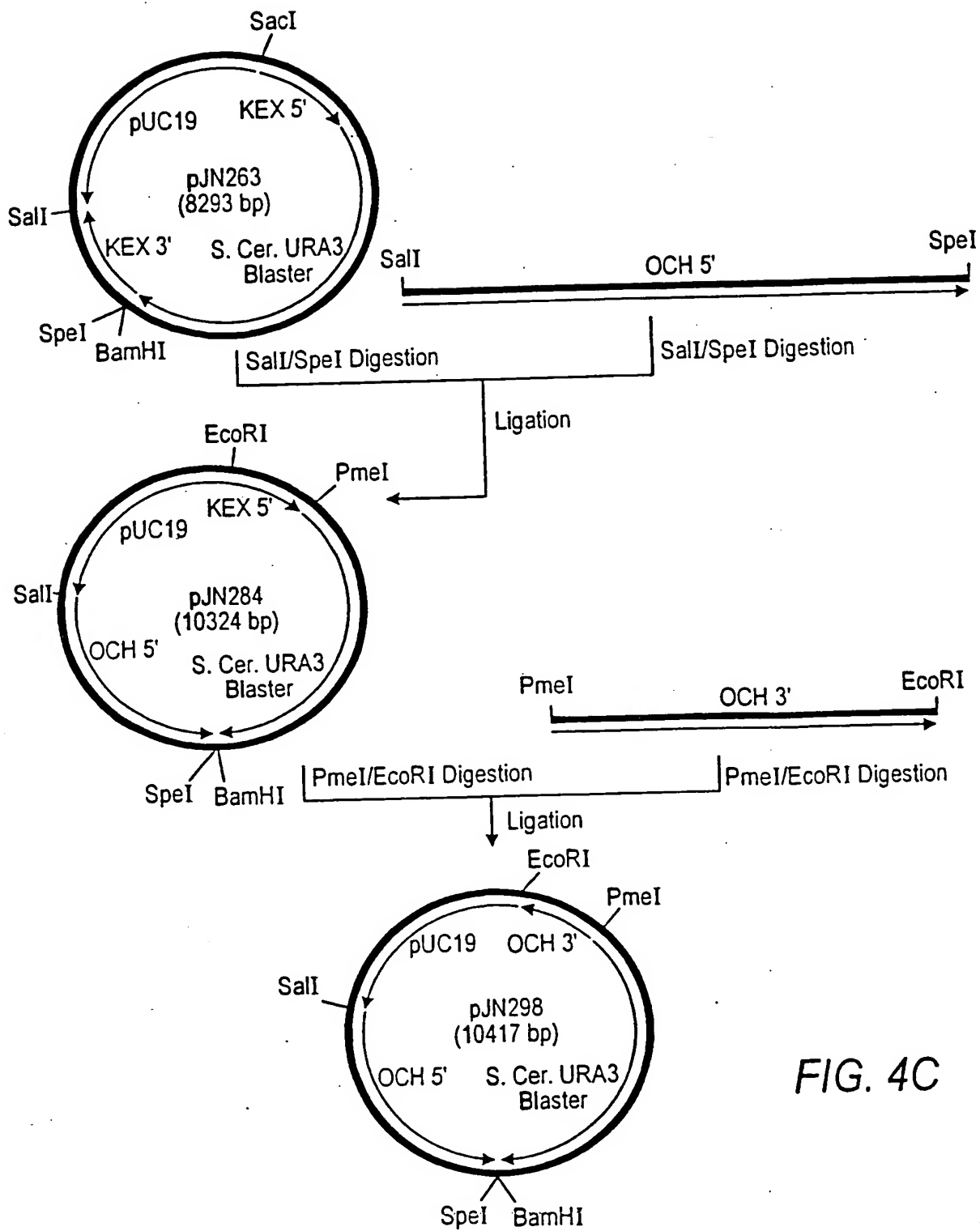


FIG. 4C



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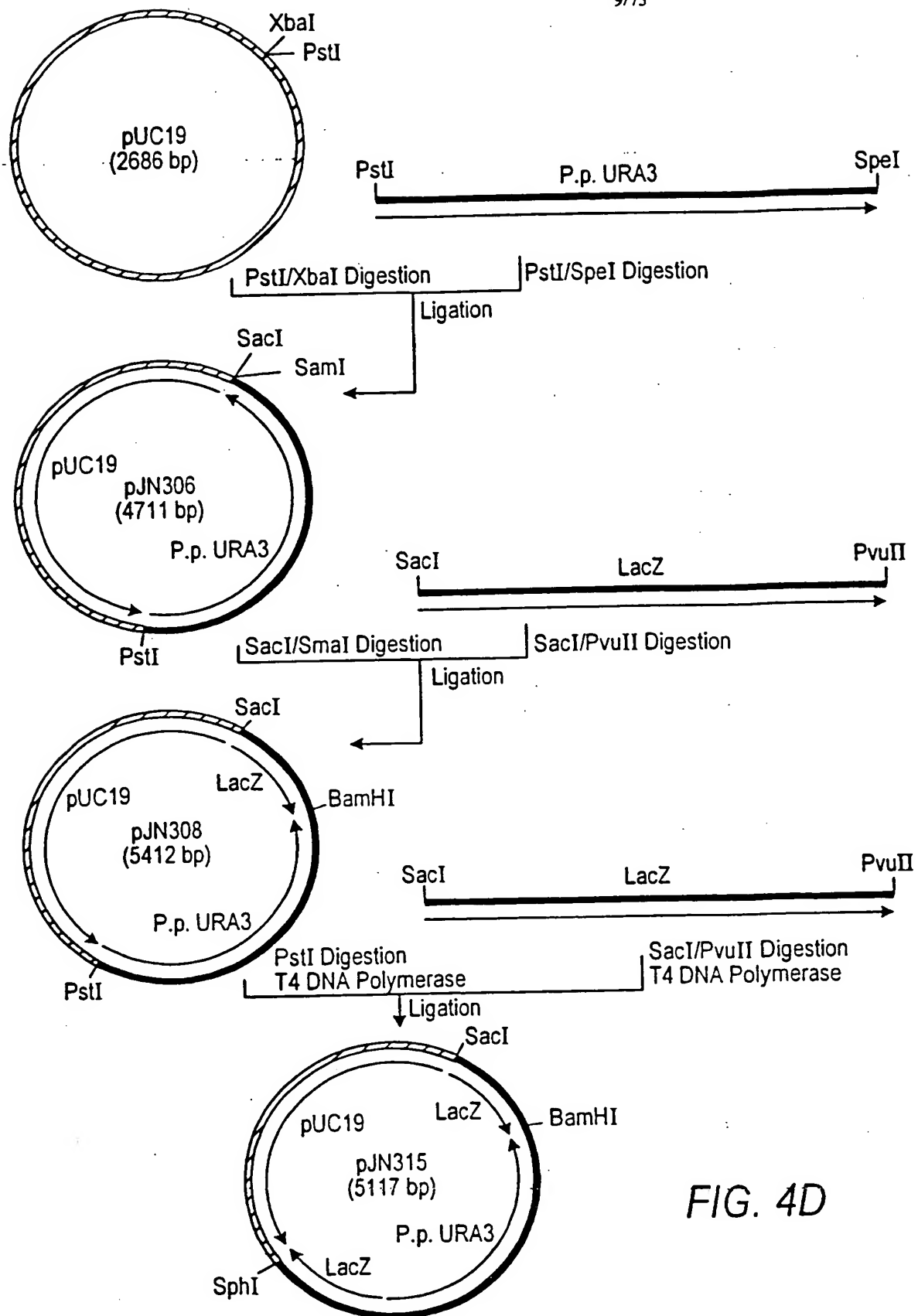


FIG. 4D

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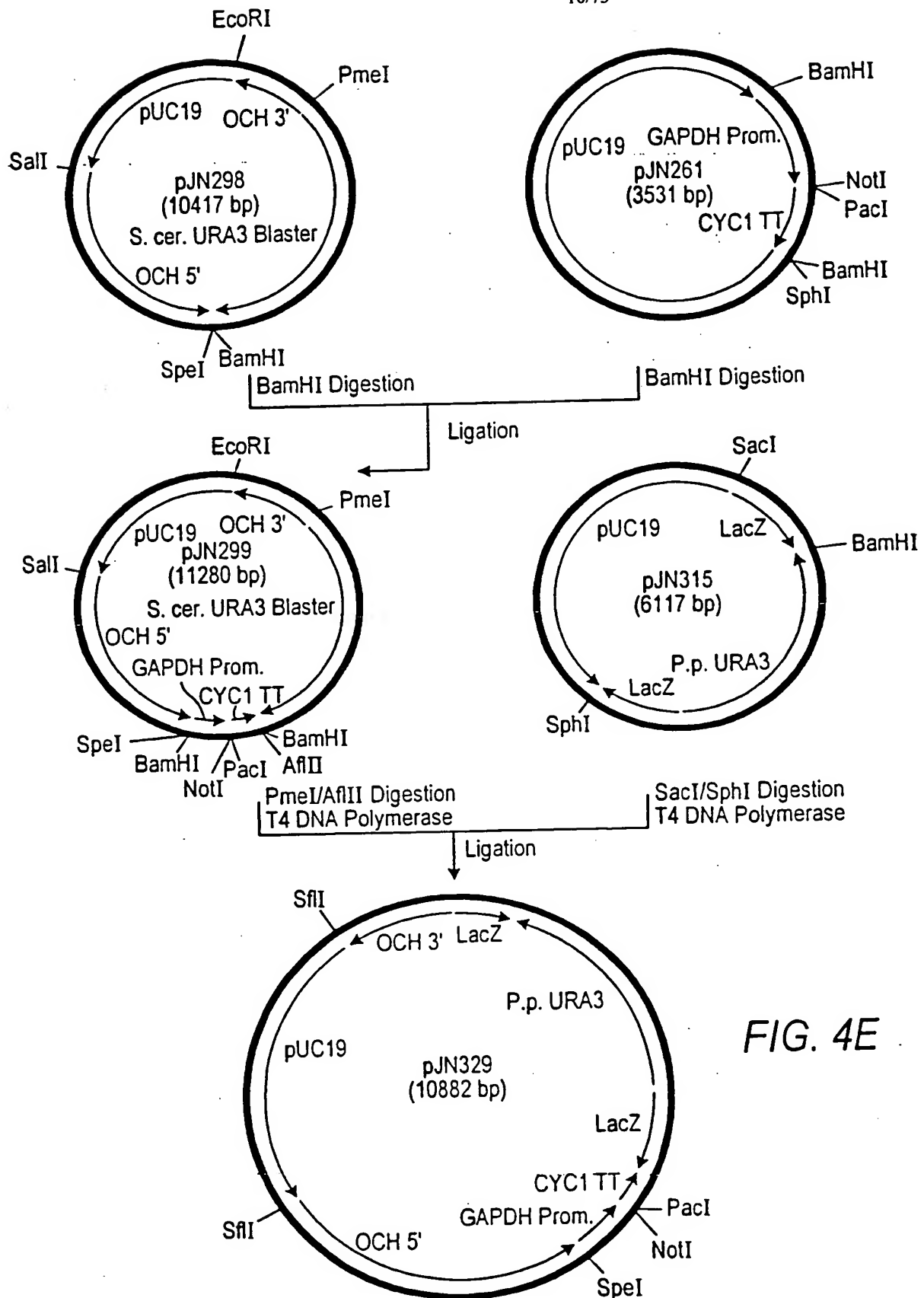


FIG. 4E

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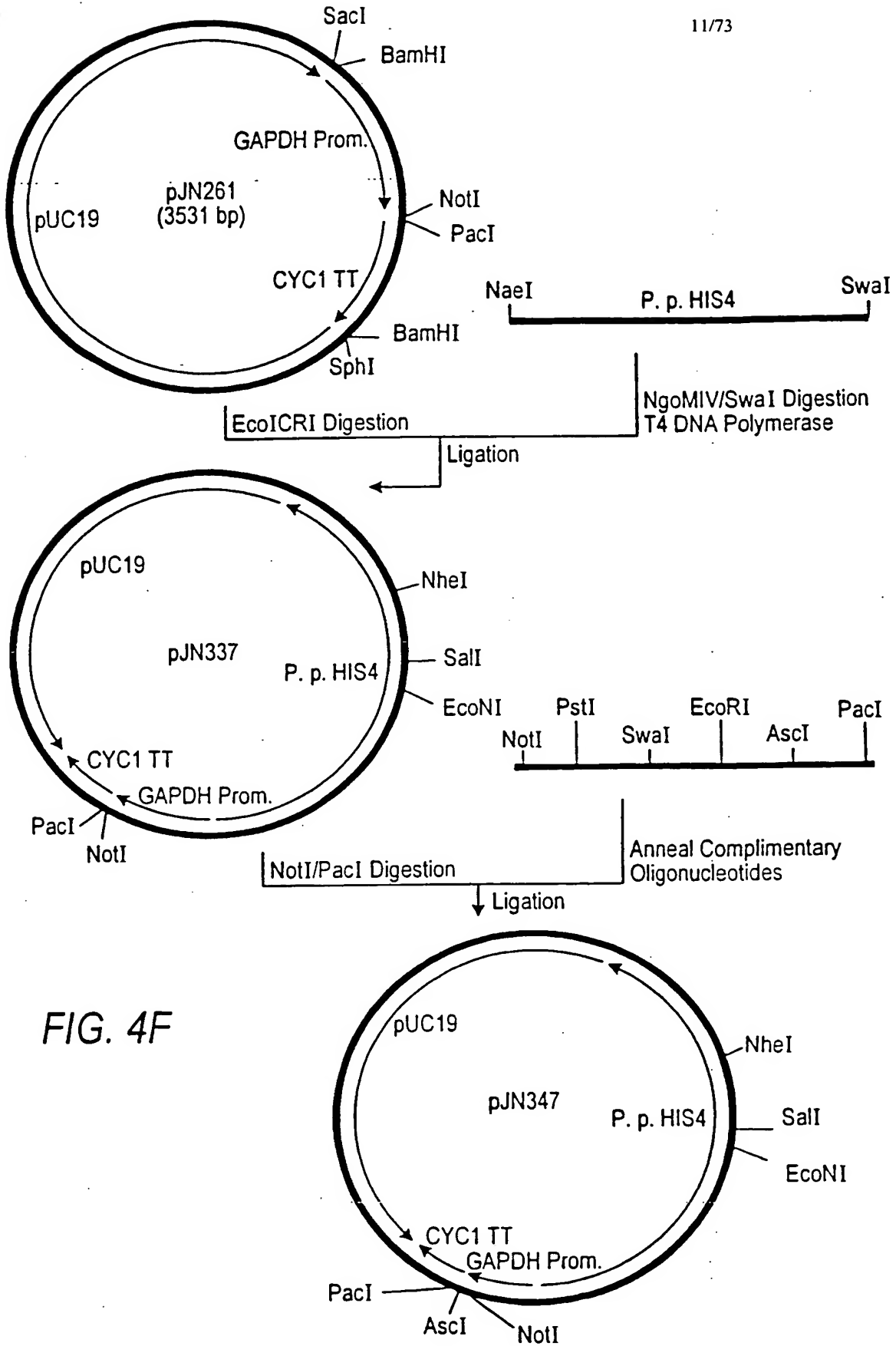


FIG. 4F

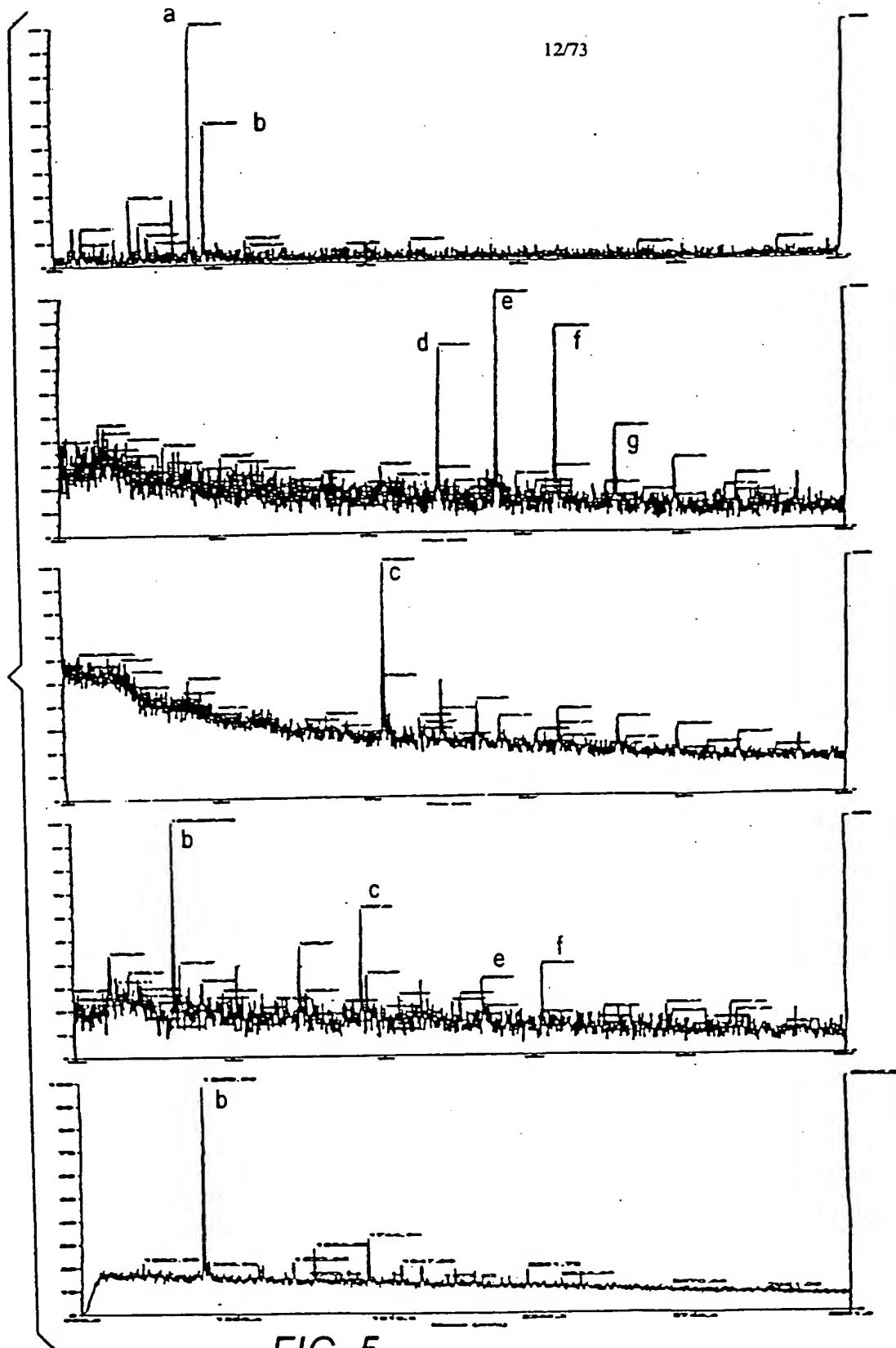
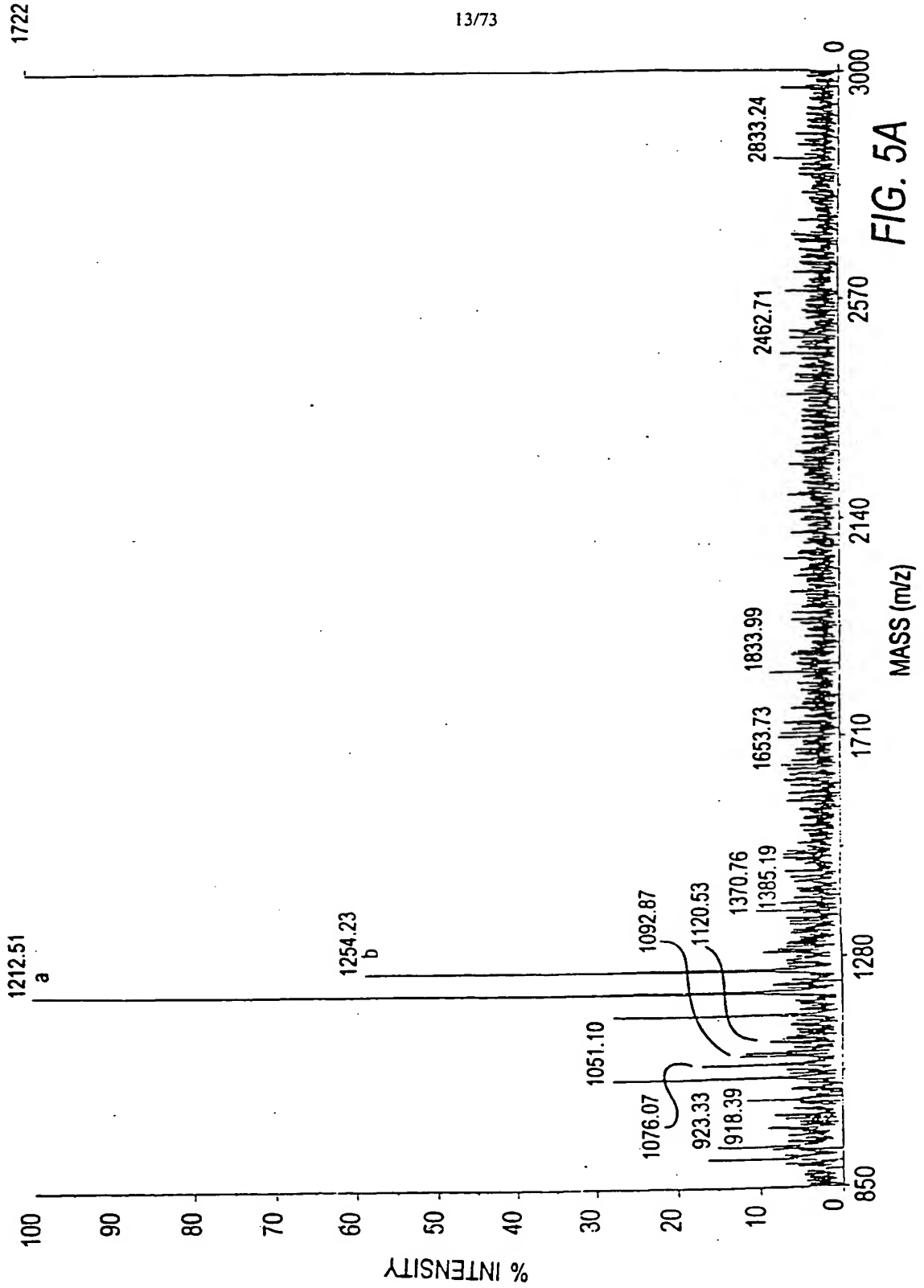


FIG. 5



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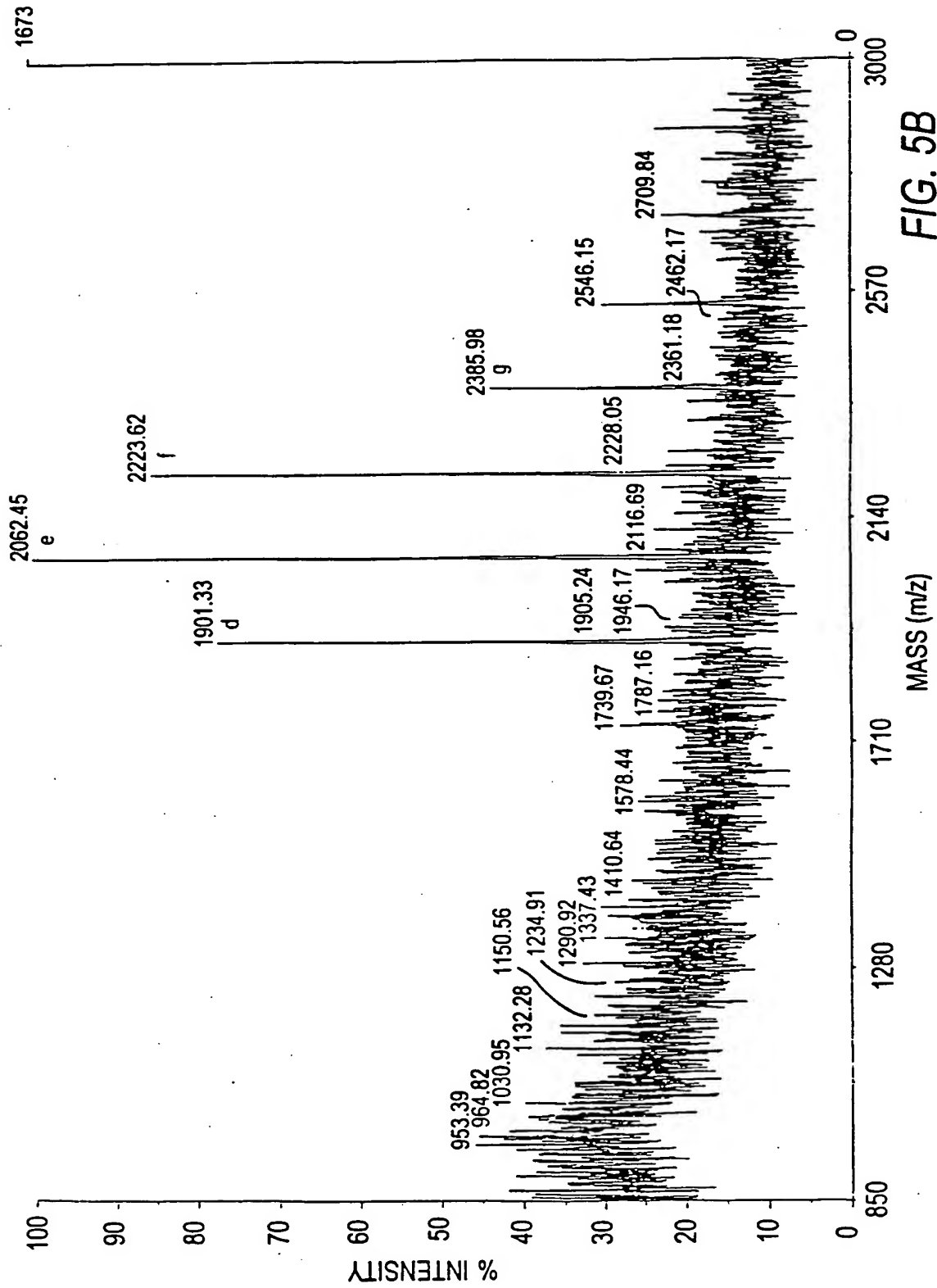


FIG. 5B

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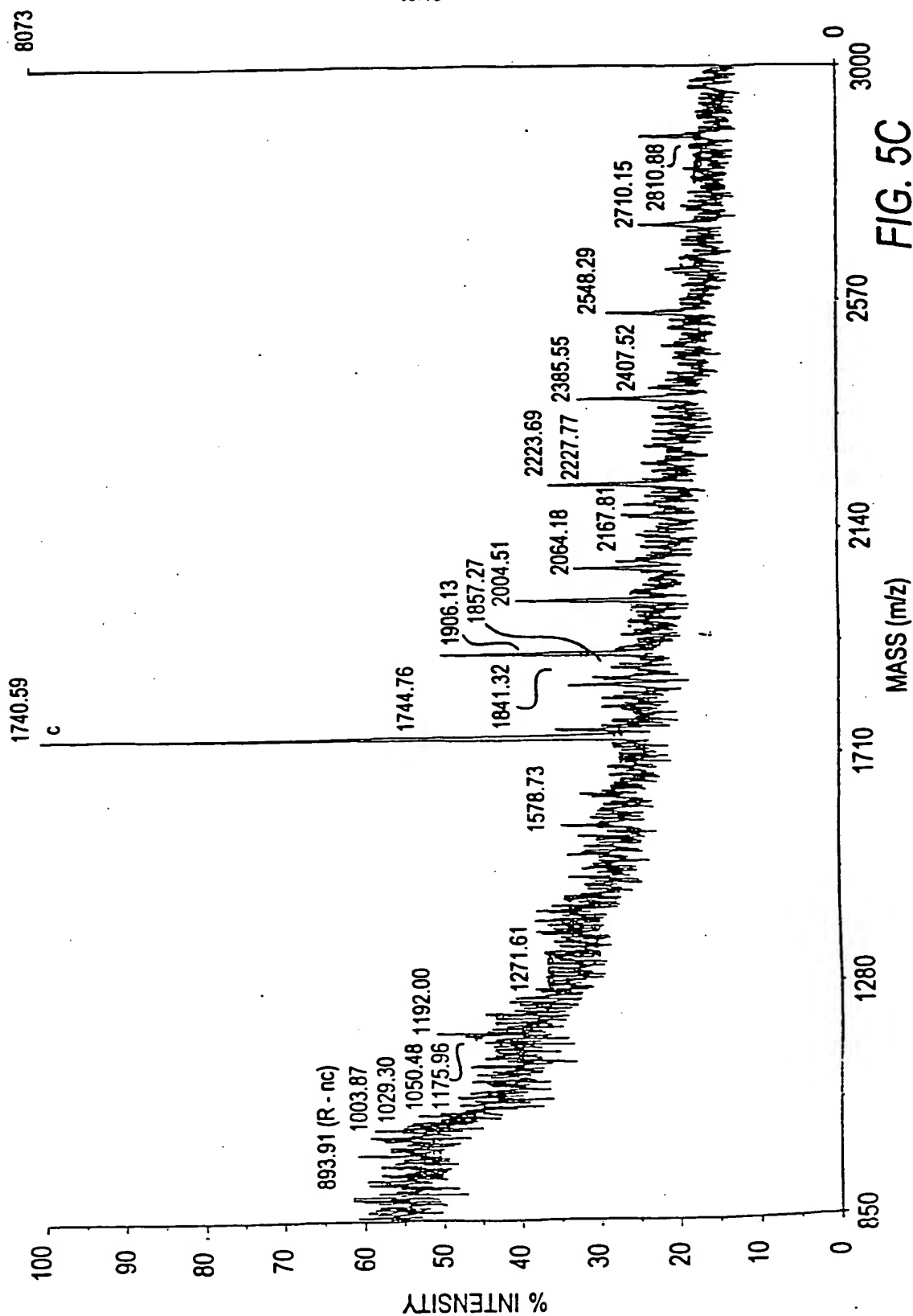


FIG. 5C

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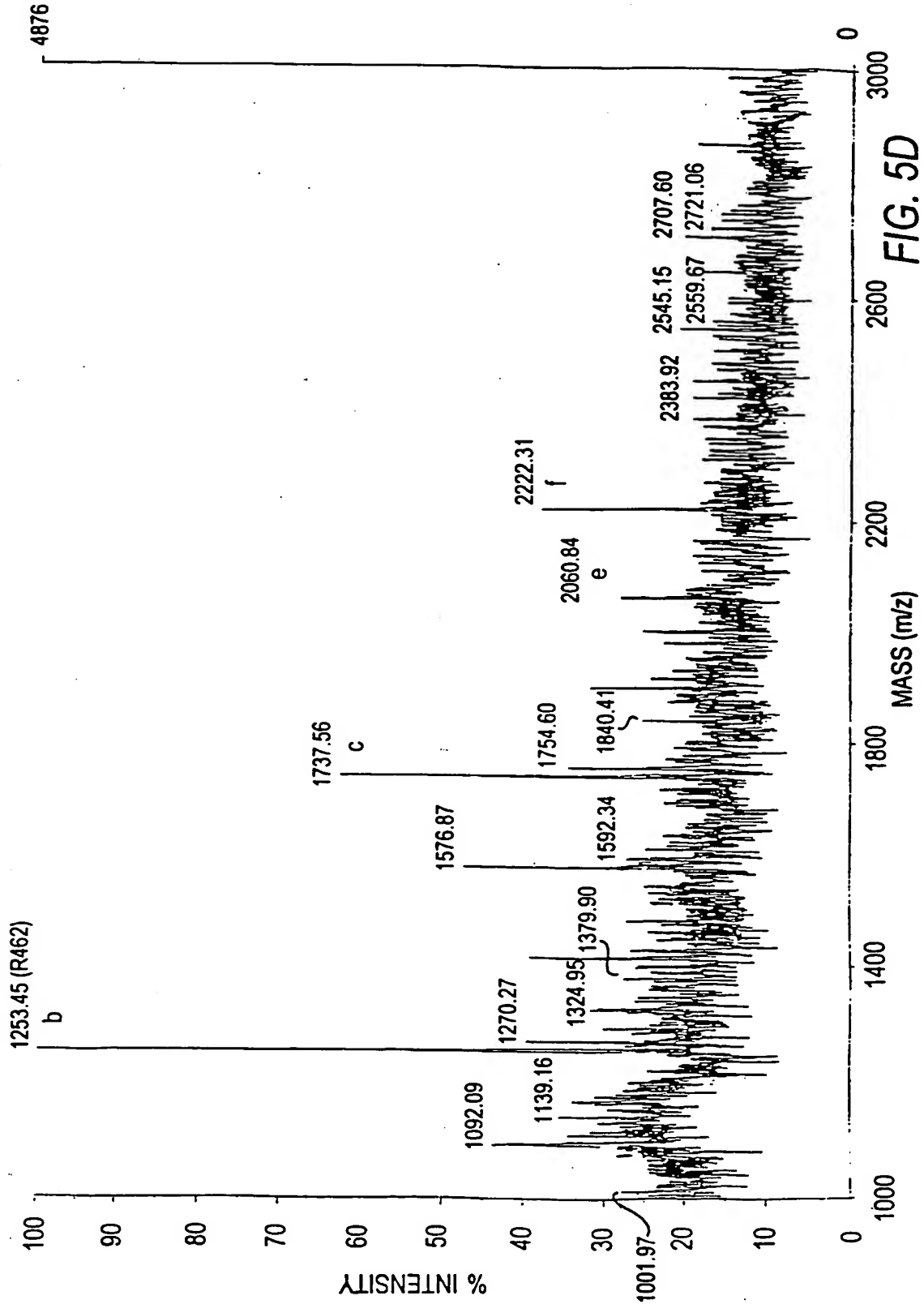


FIG. 5D



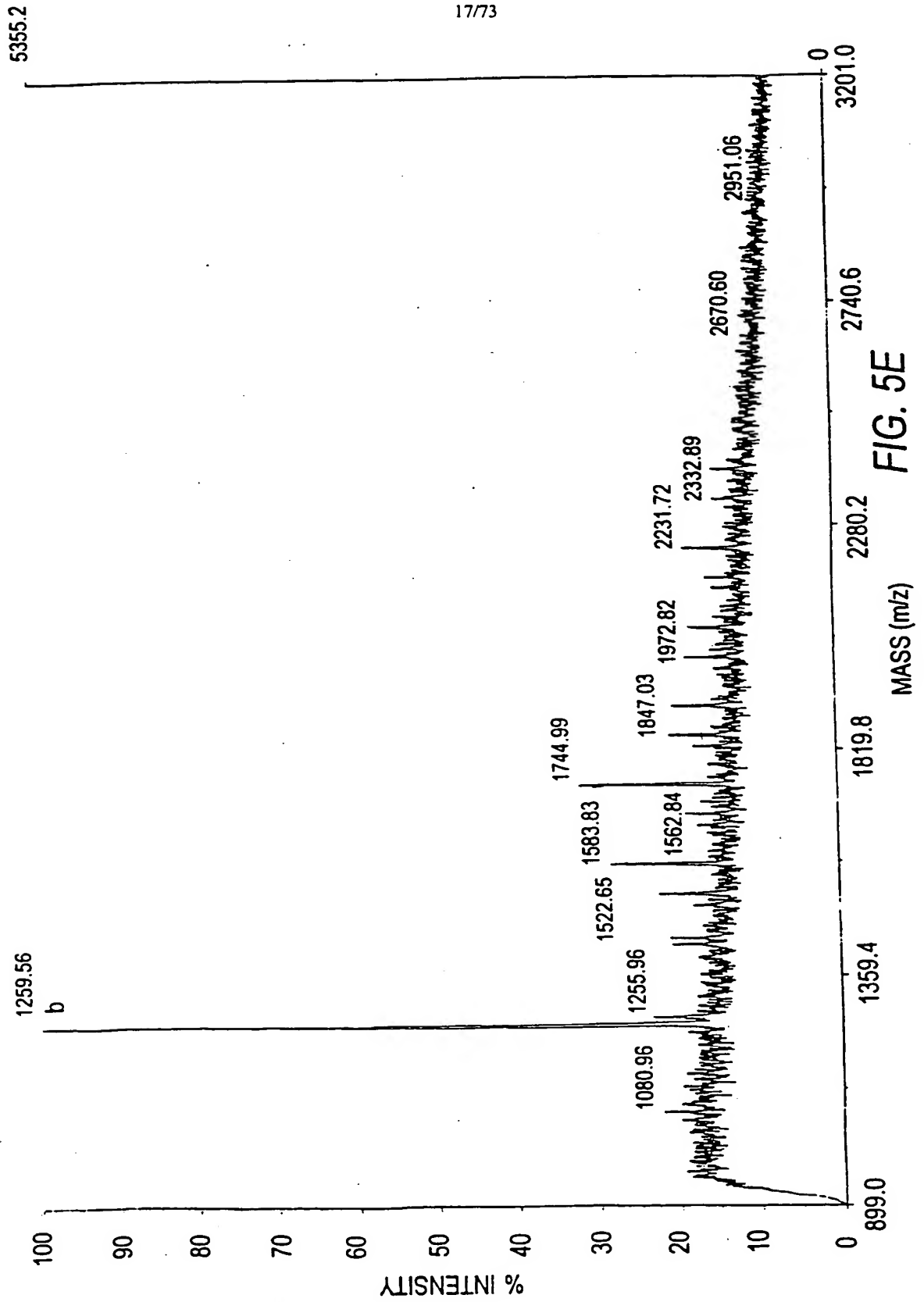
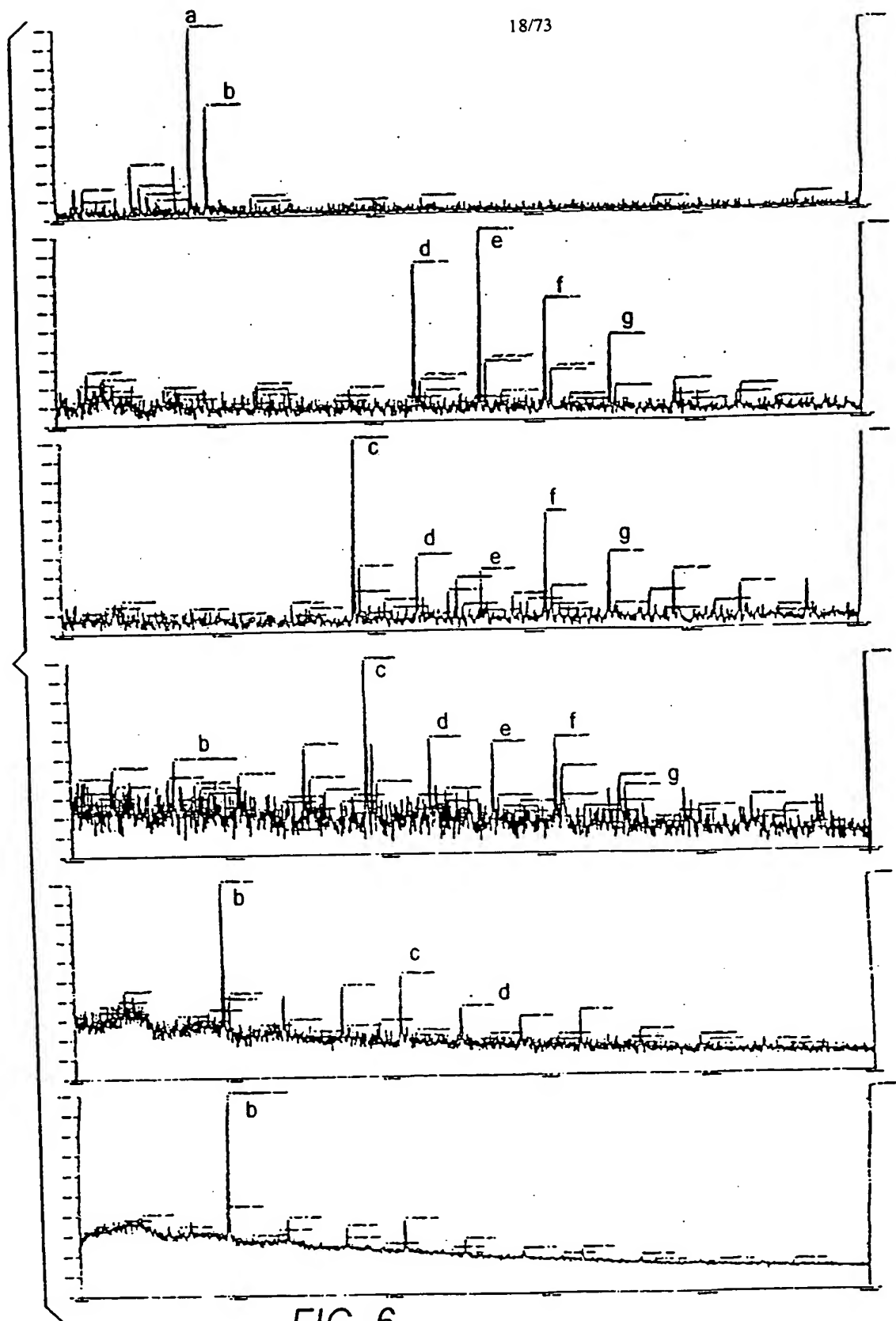
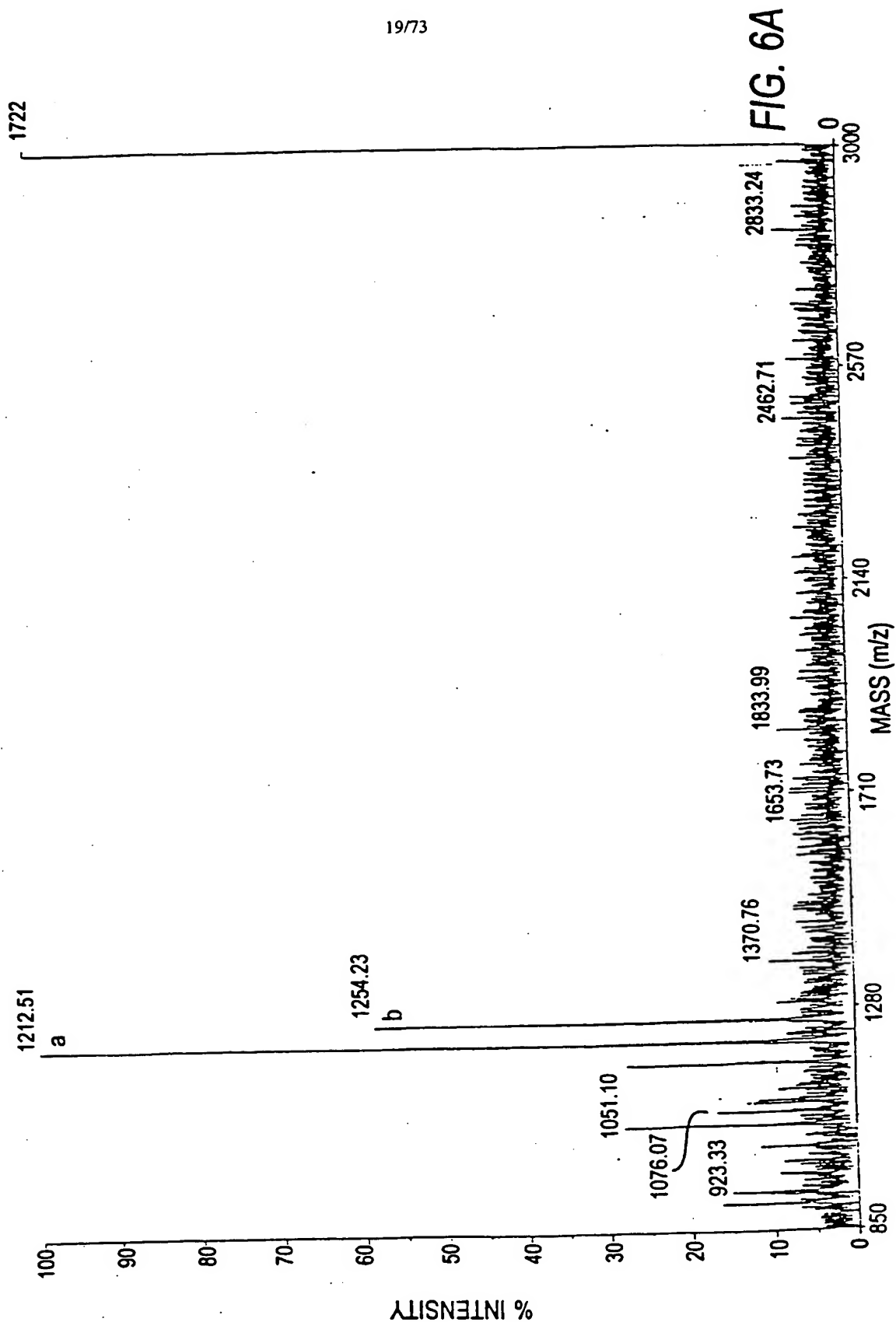


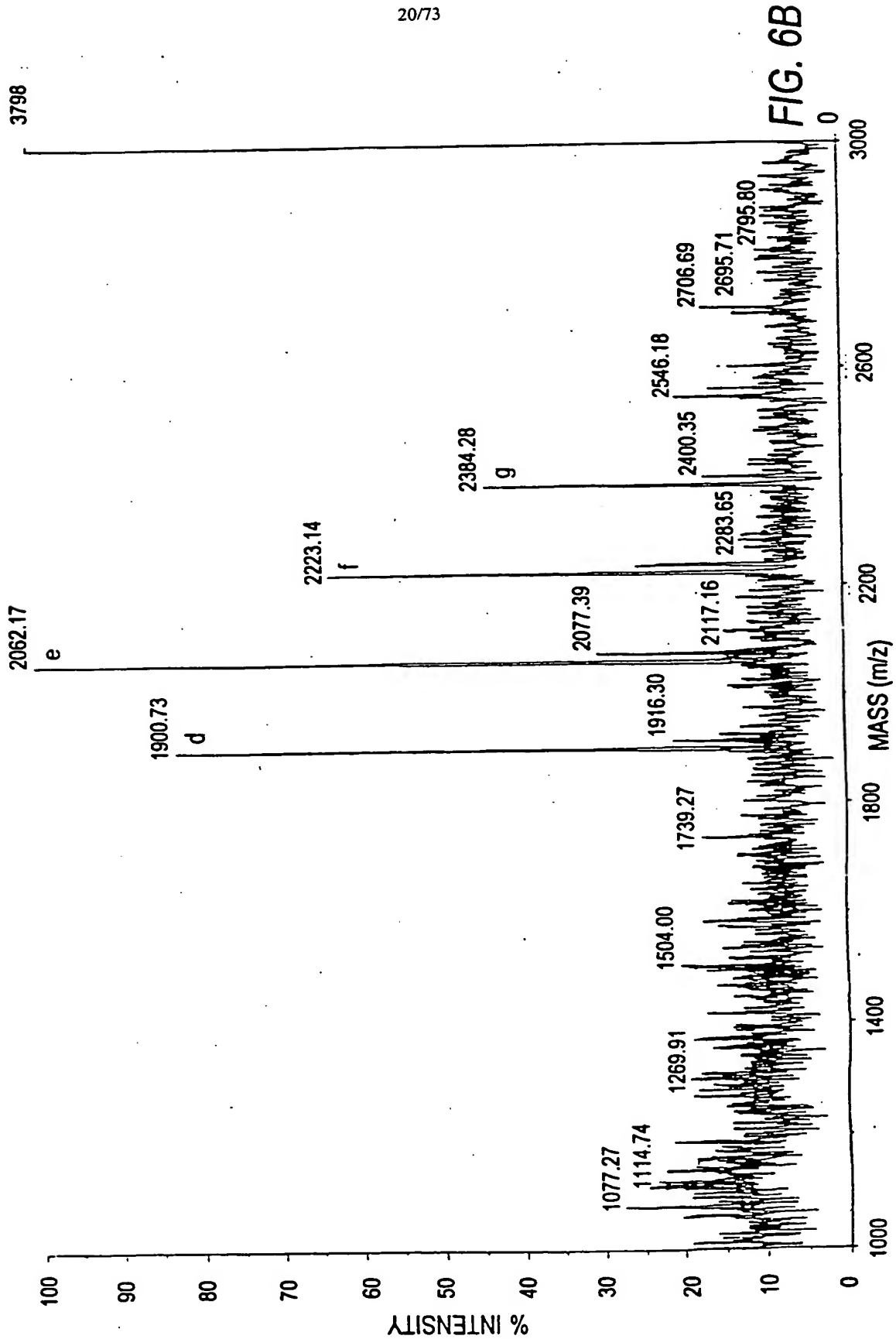
FIG. 5E



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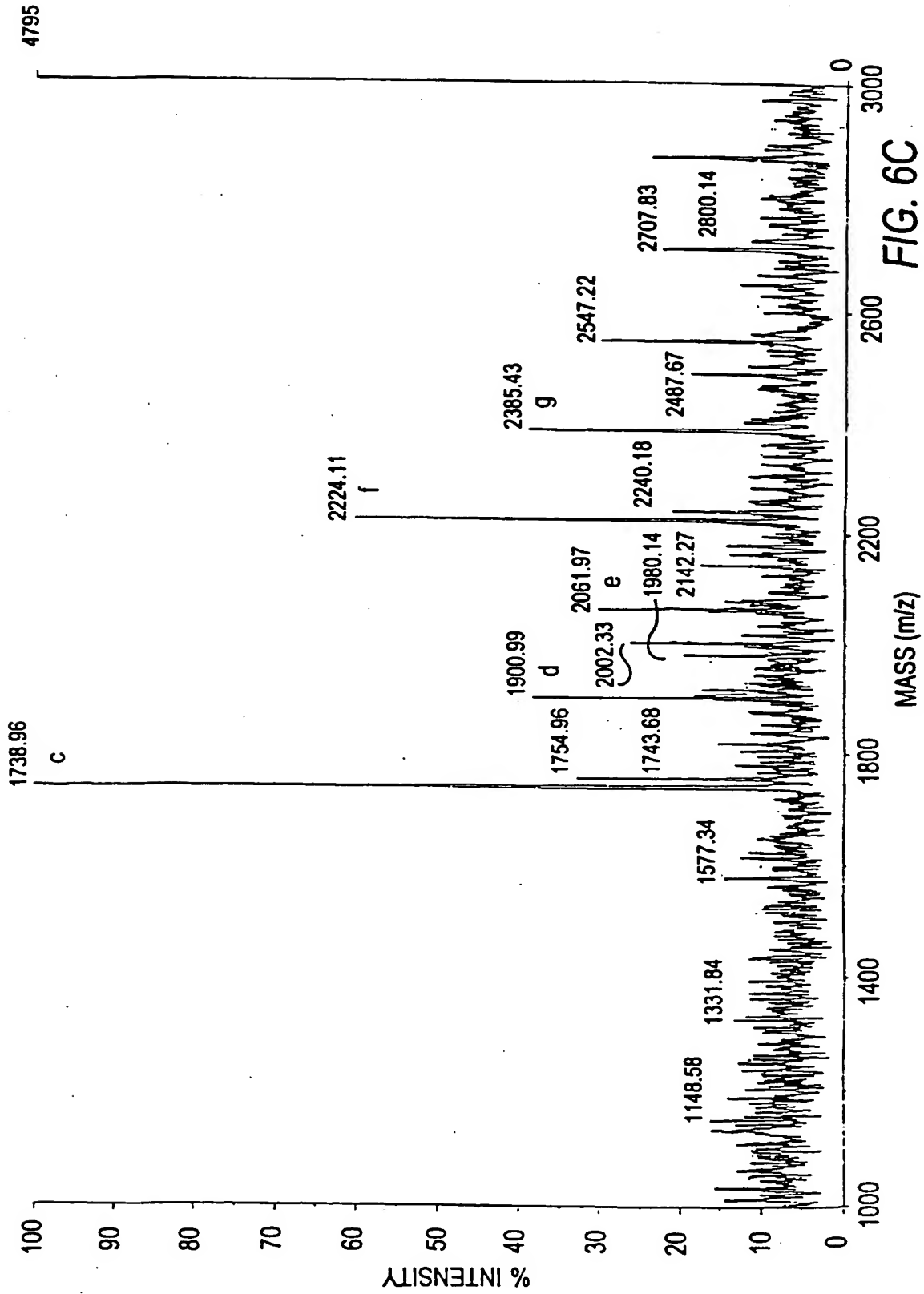
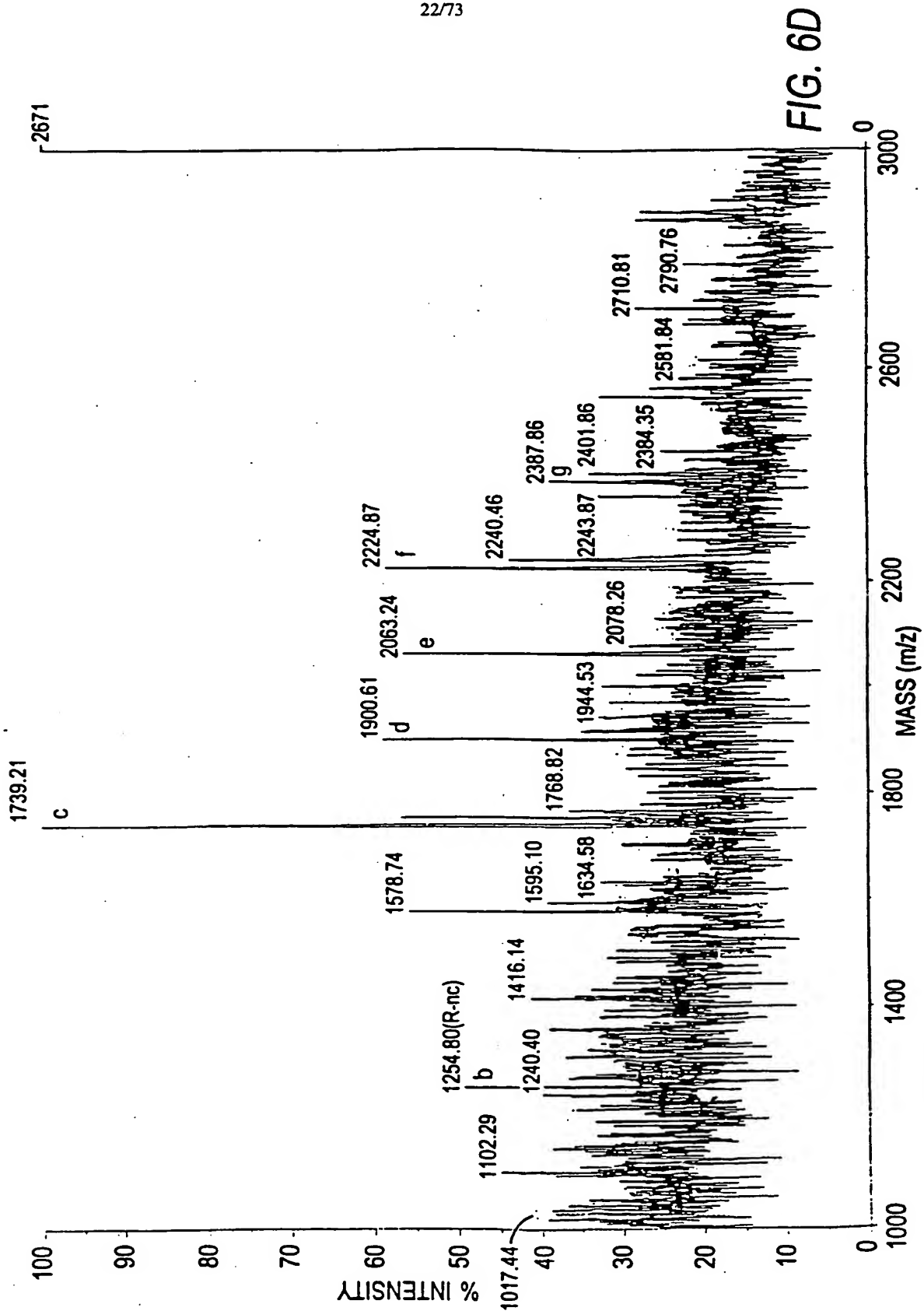


FIG. 6C

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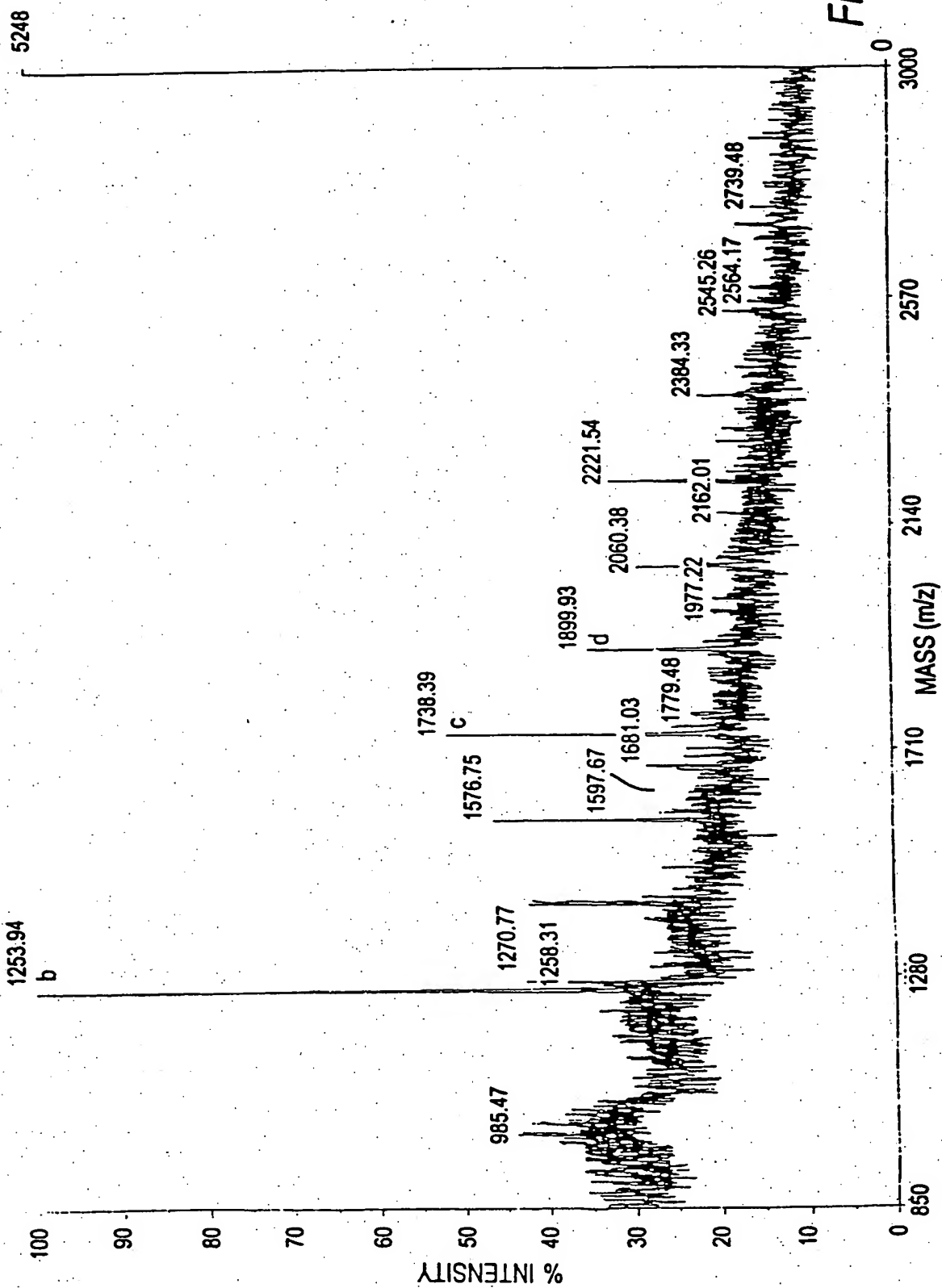


FIG. 6E

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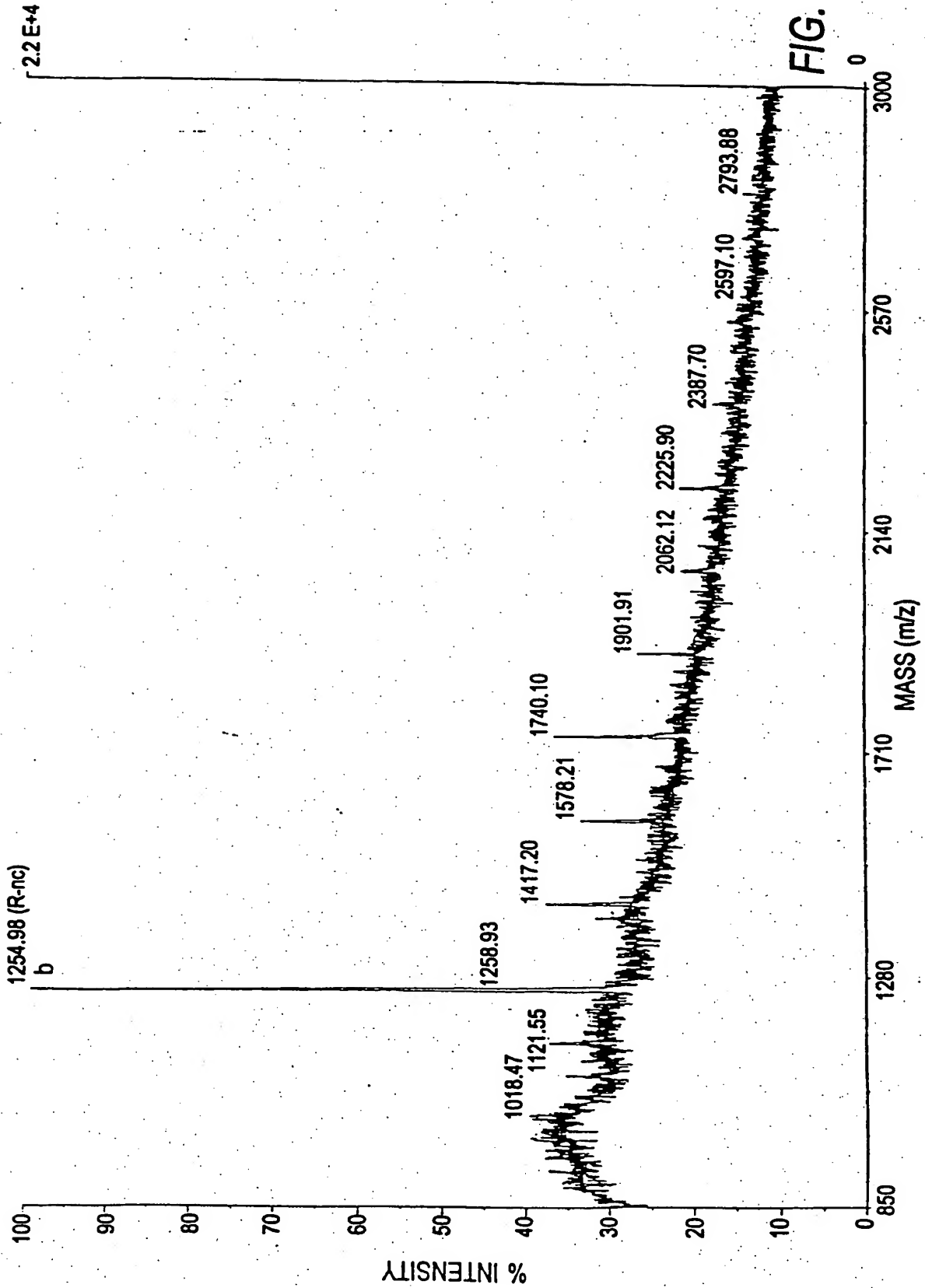


FIG. 6F



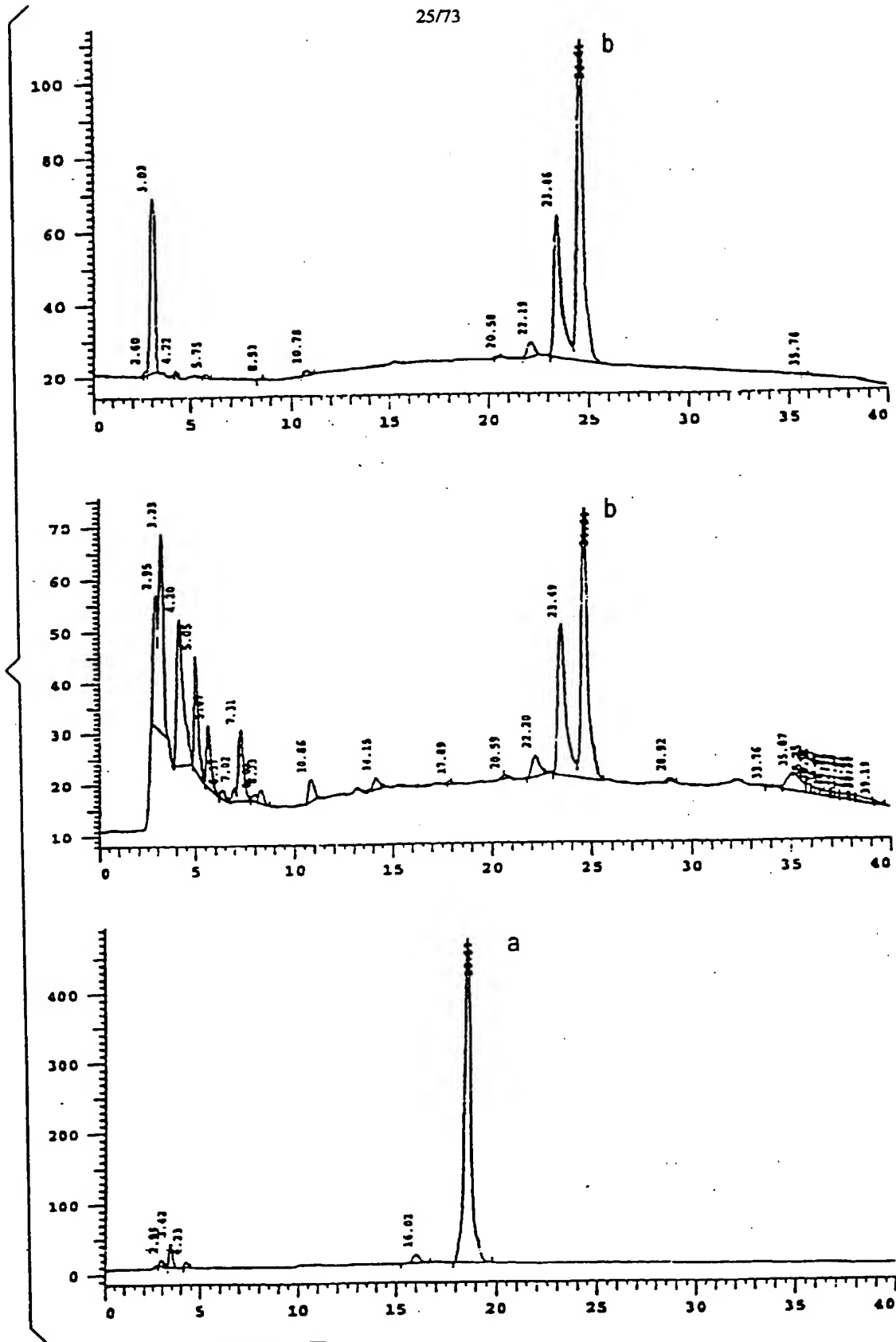


FIG. 7

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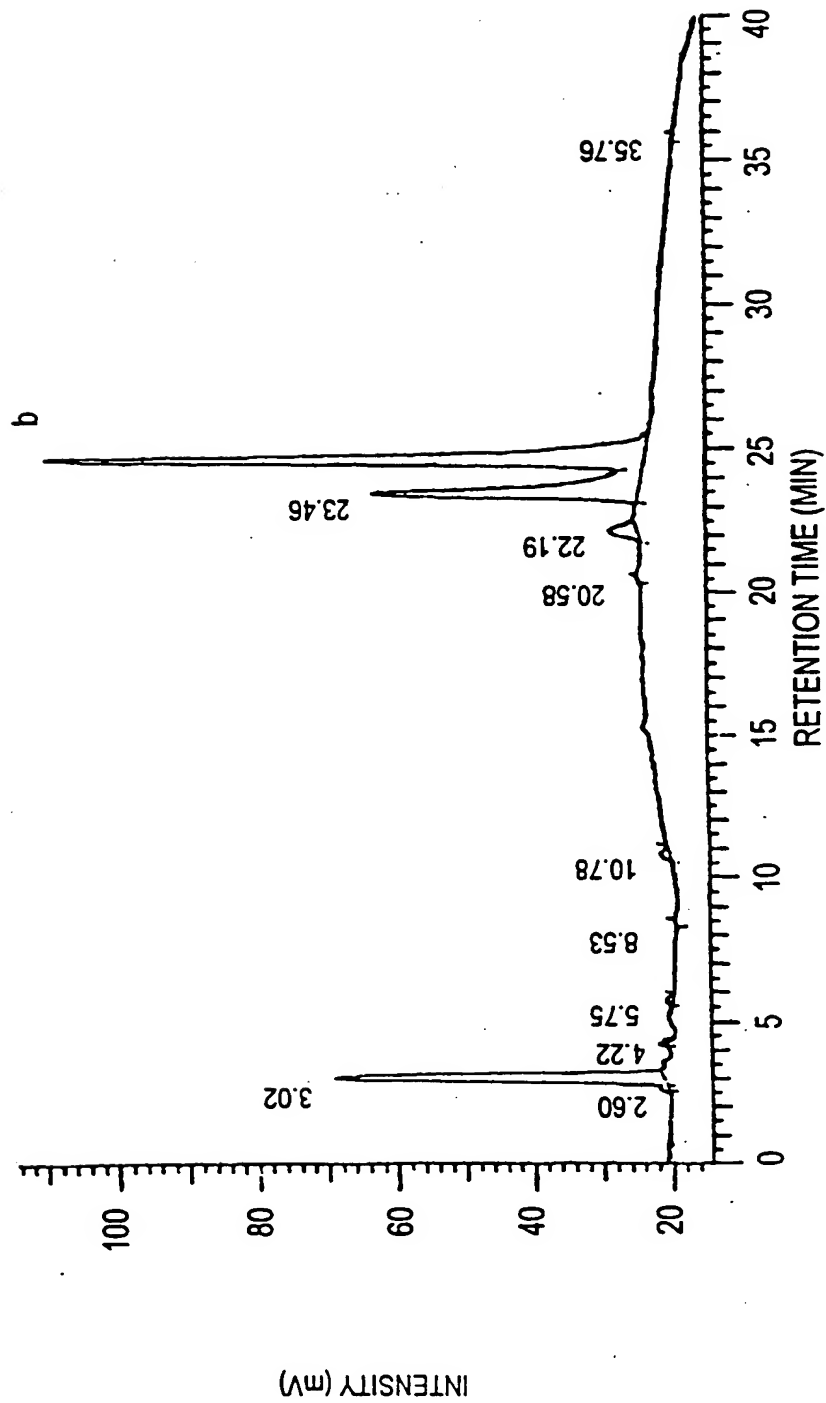


FIG. 7A

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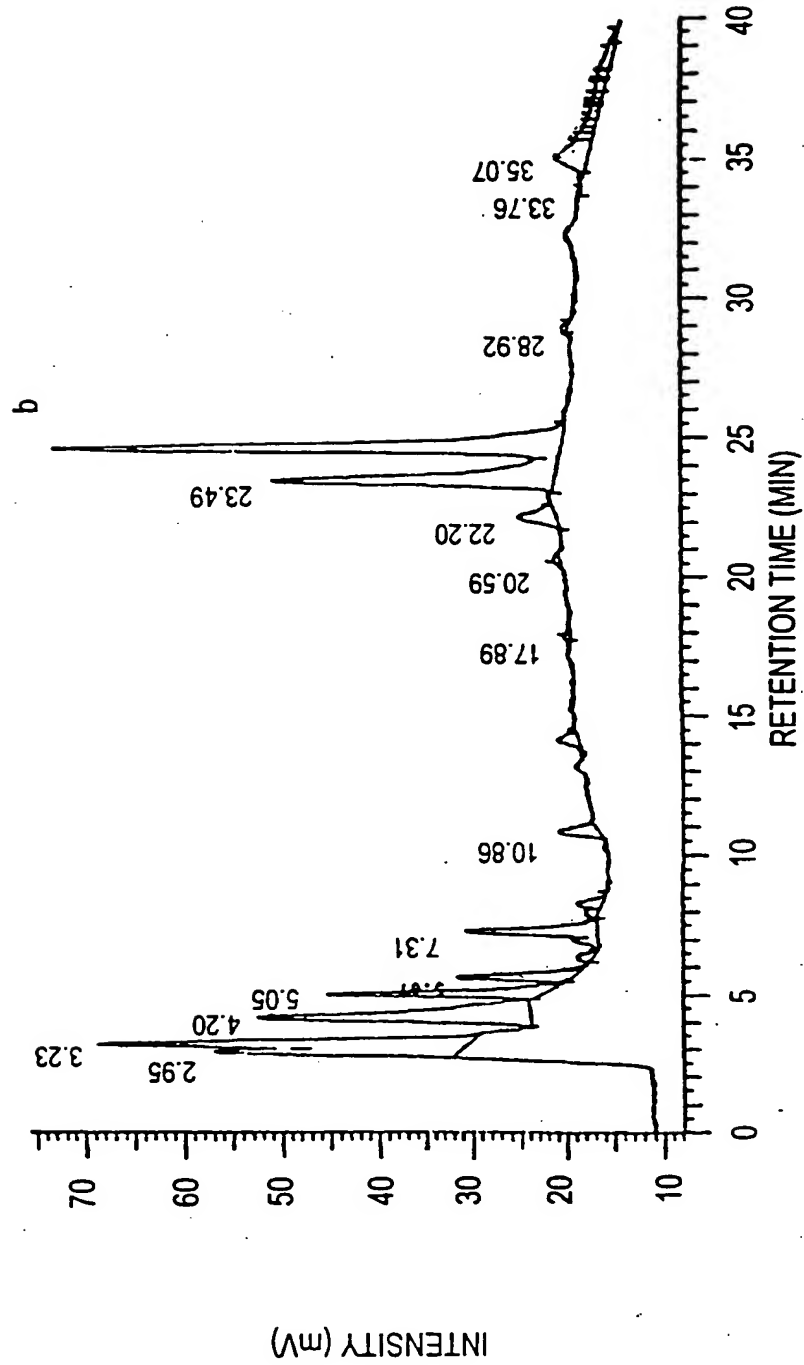


FIG. 7B

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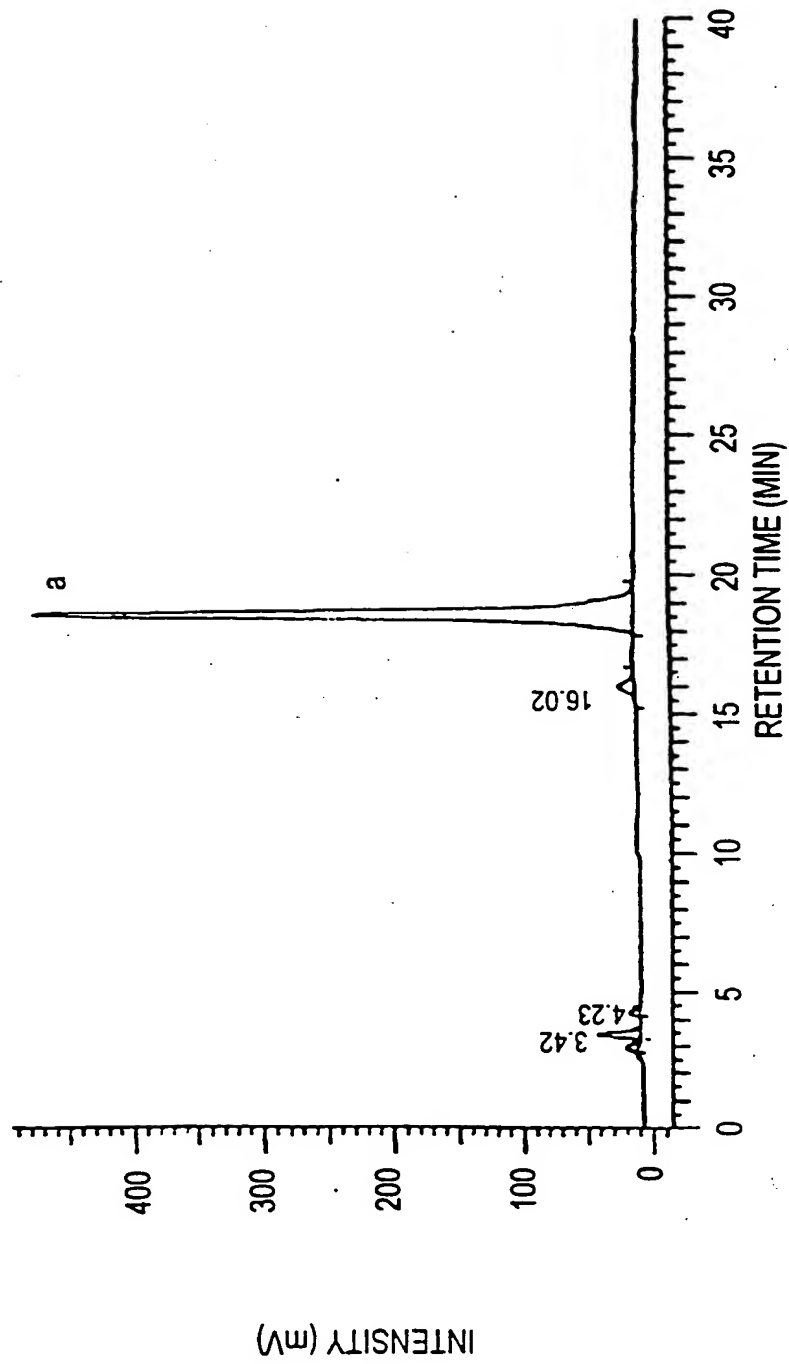


FIG. 7C

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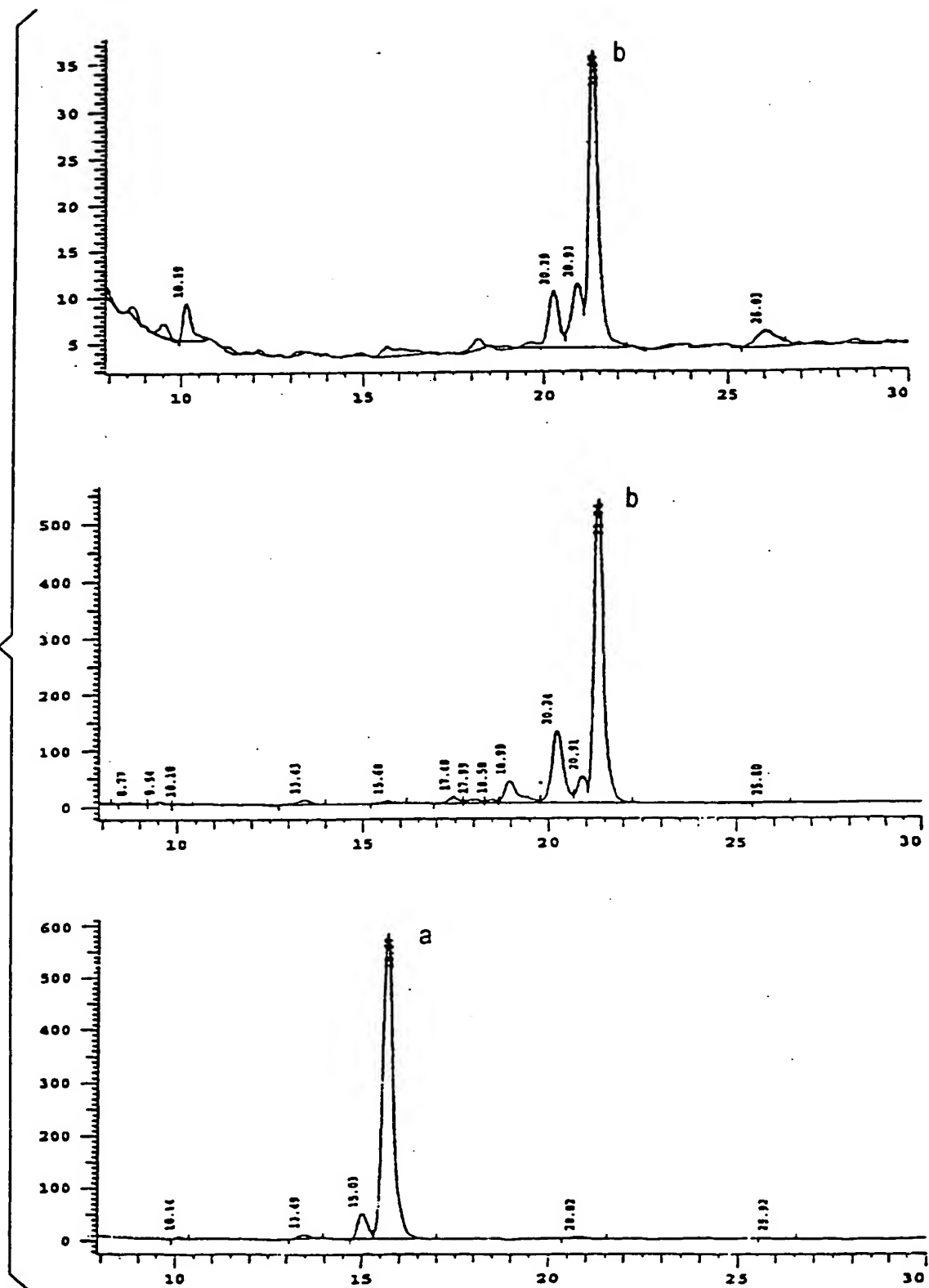


FIG. 8

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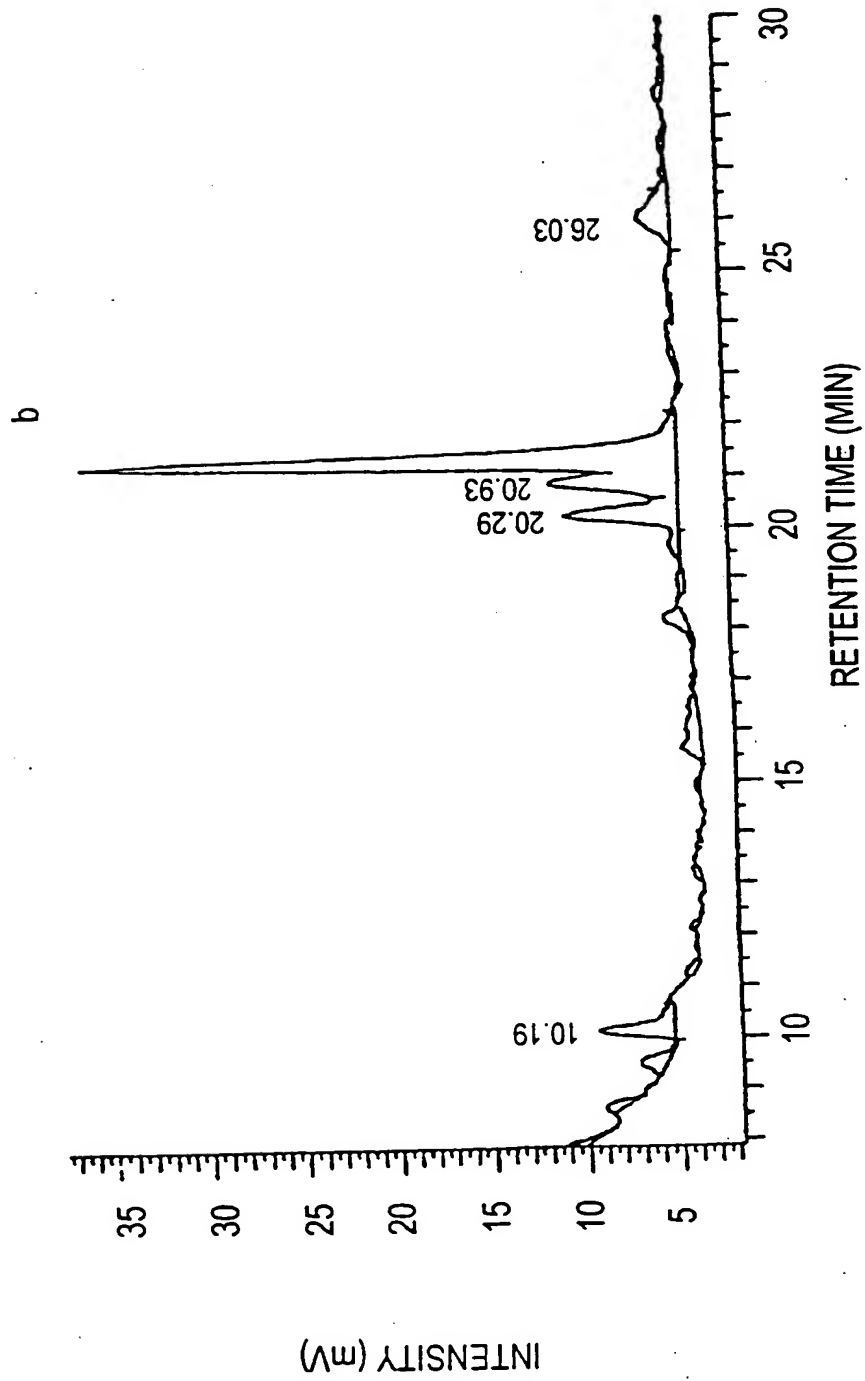


FIG. 8A

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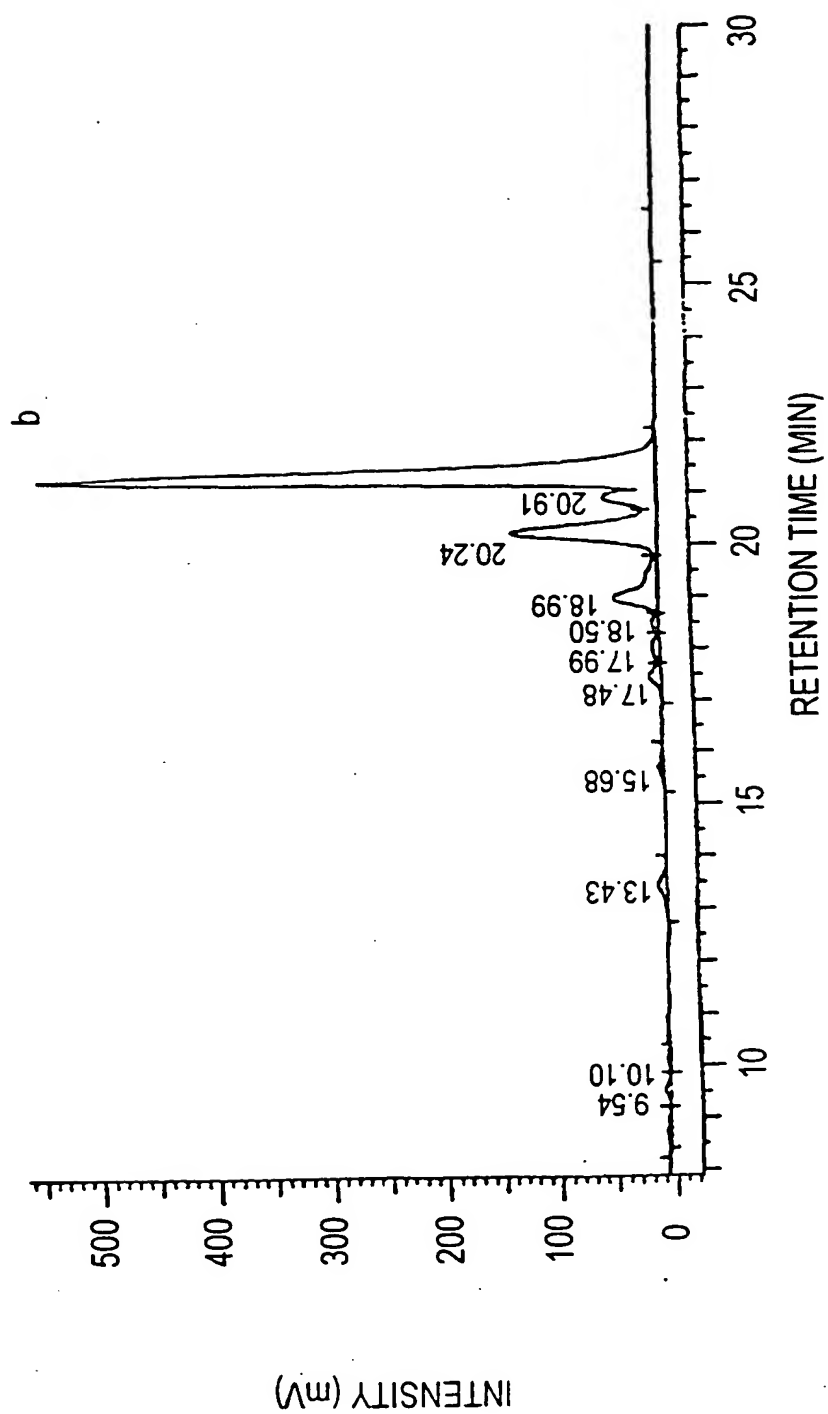


FIG. 8B

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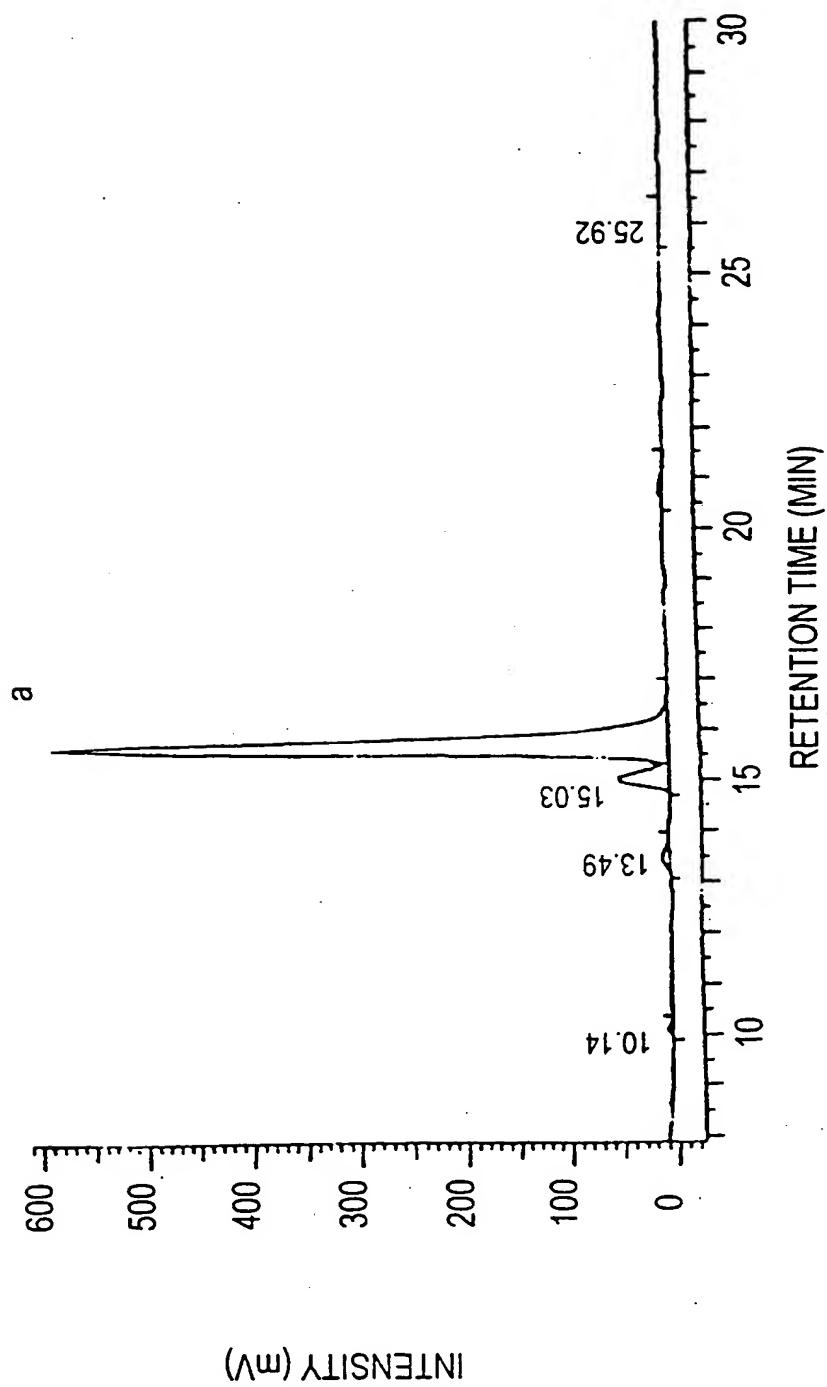


FIG. 8C



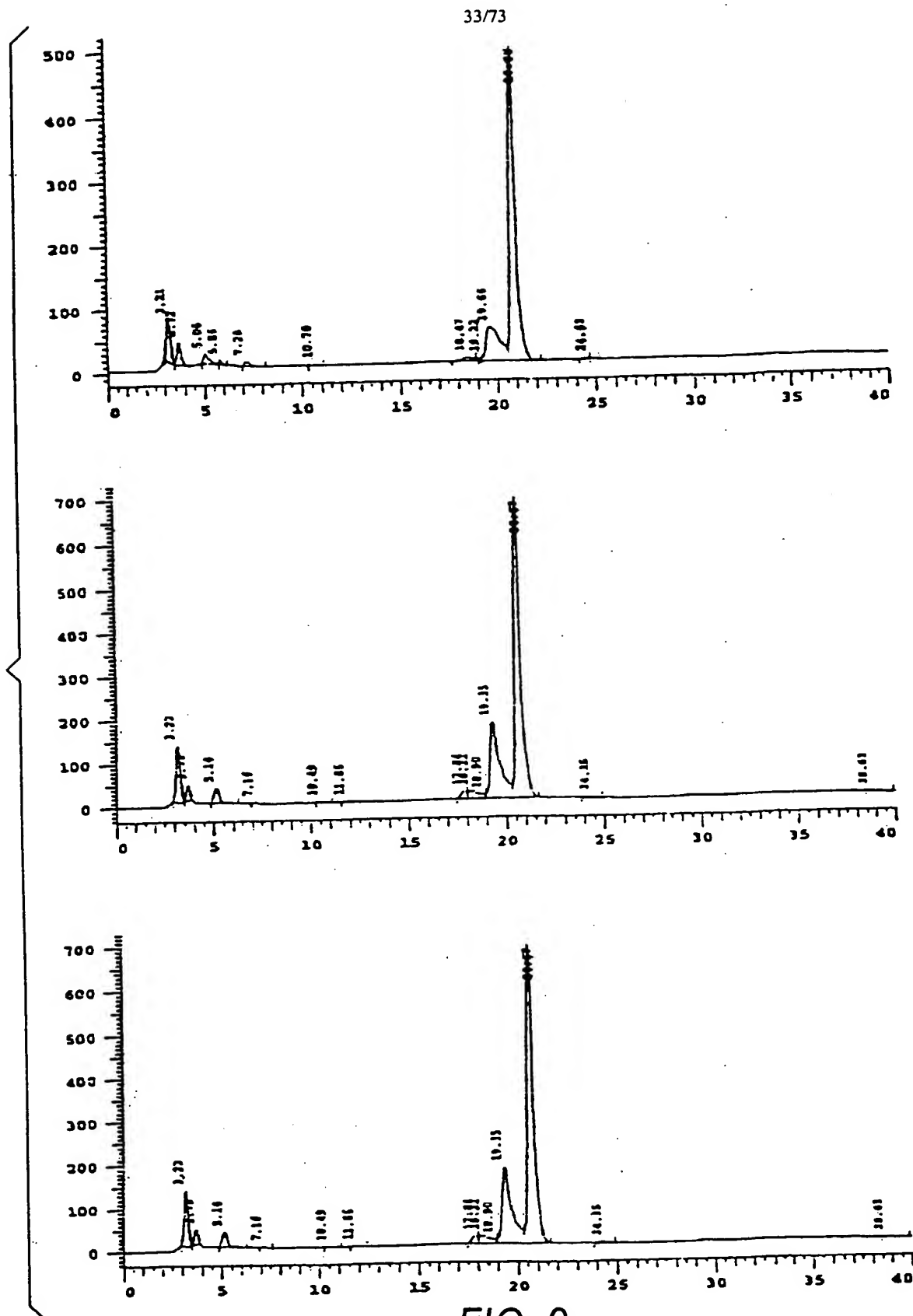


FIG. 9

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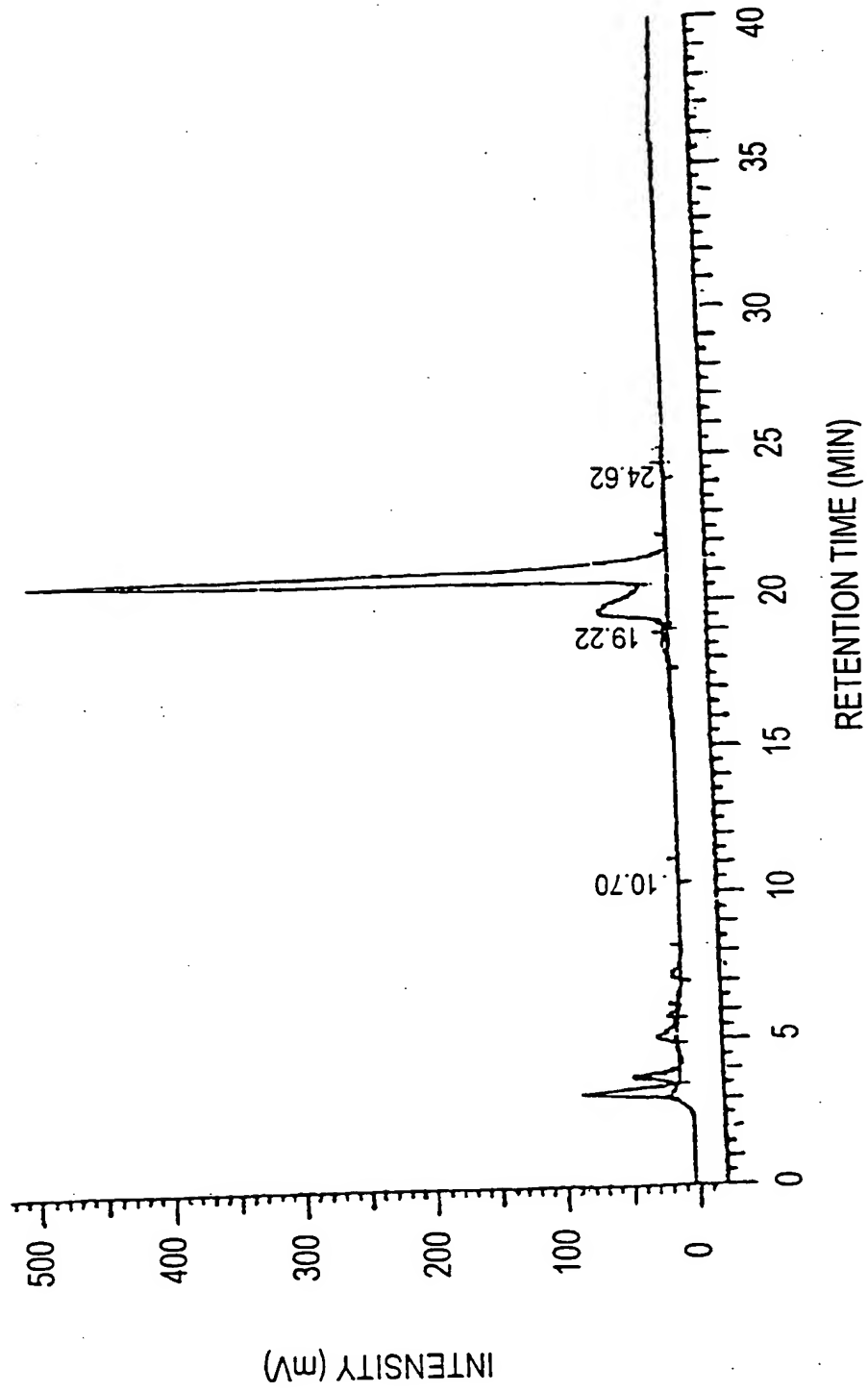


FIG. 9A

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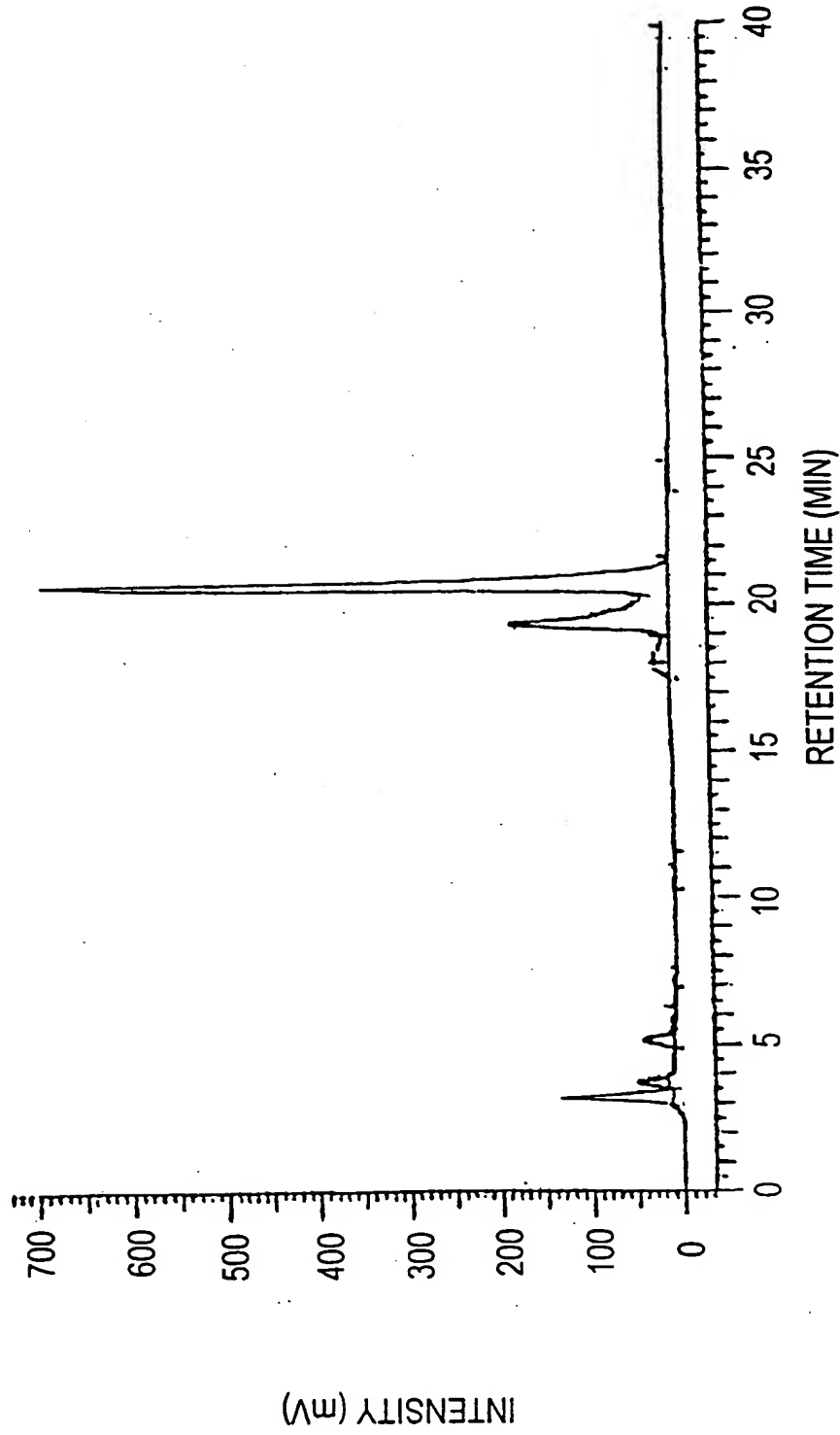


FIG. 9B

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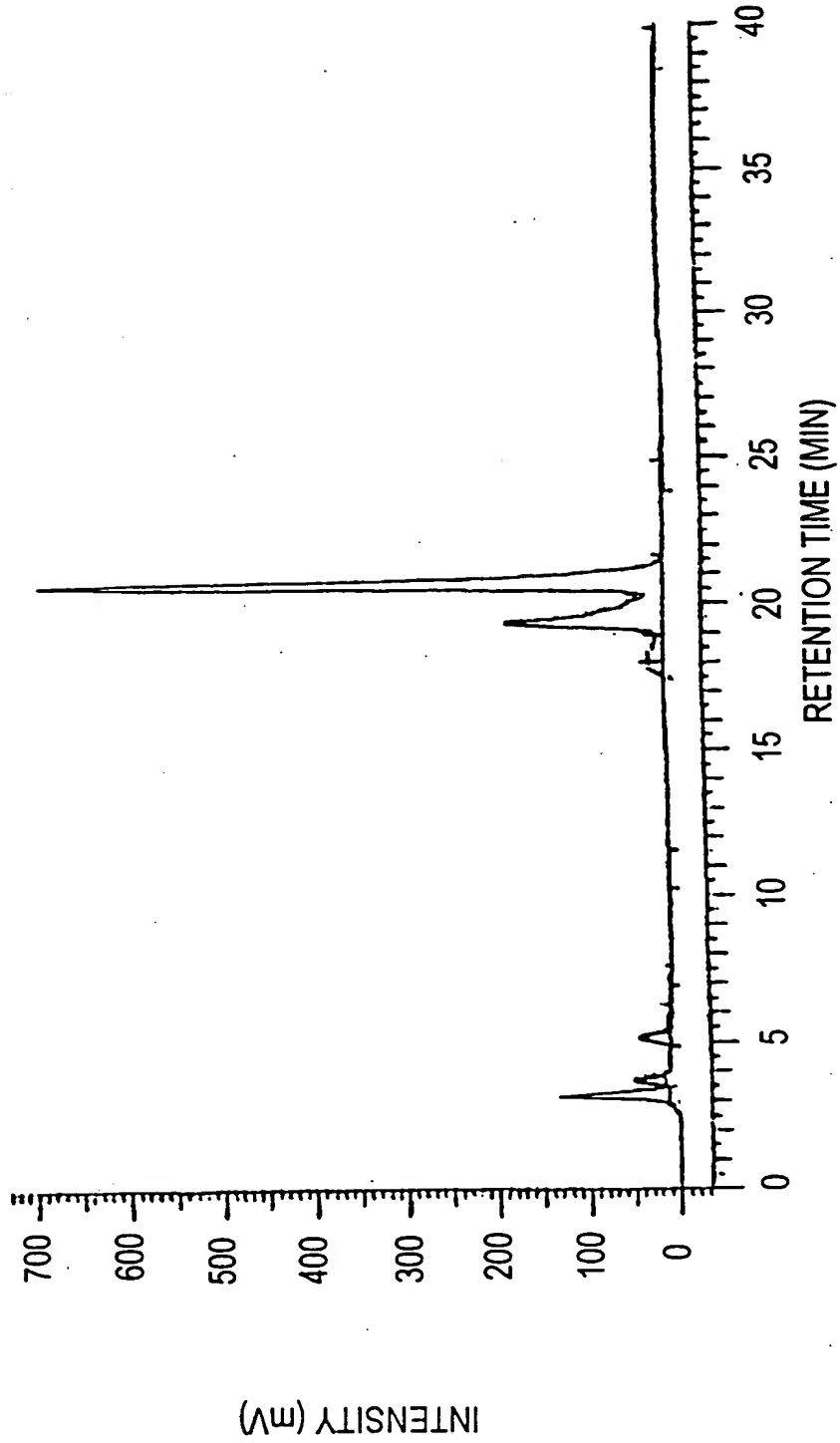
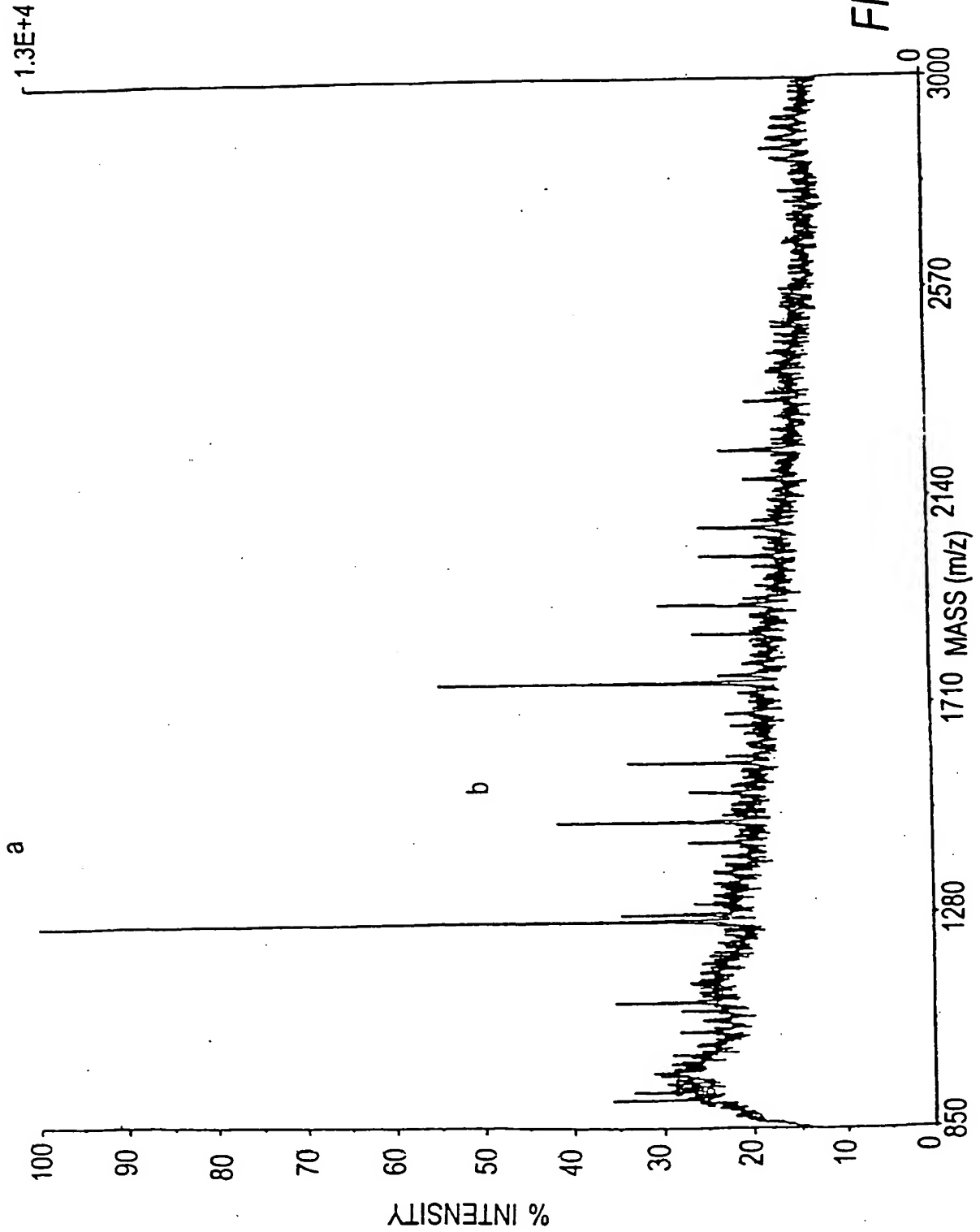


FIG. 9C



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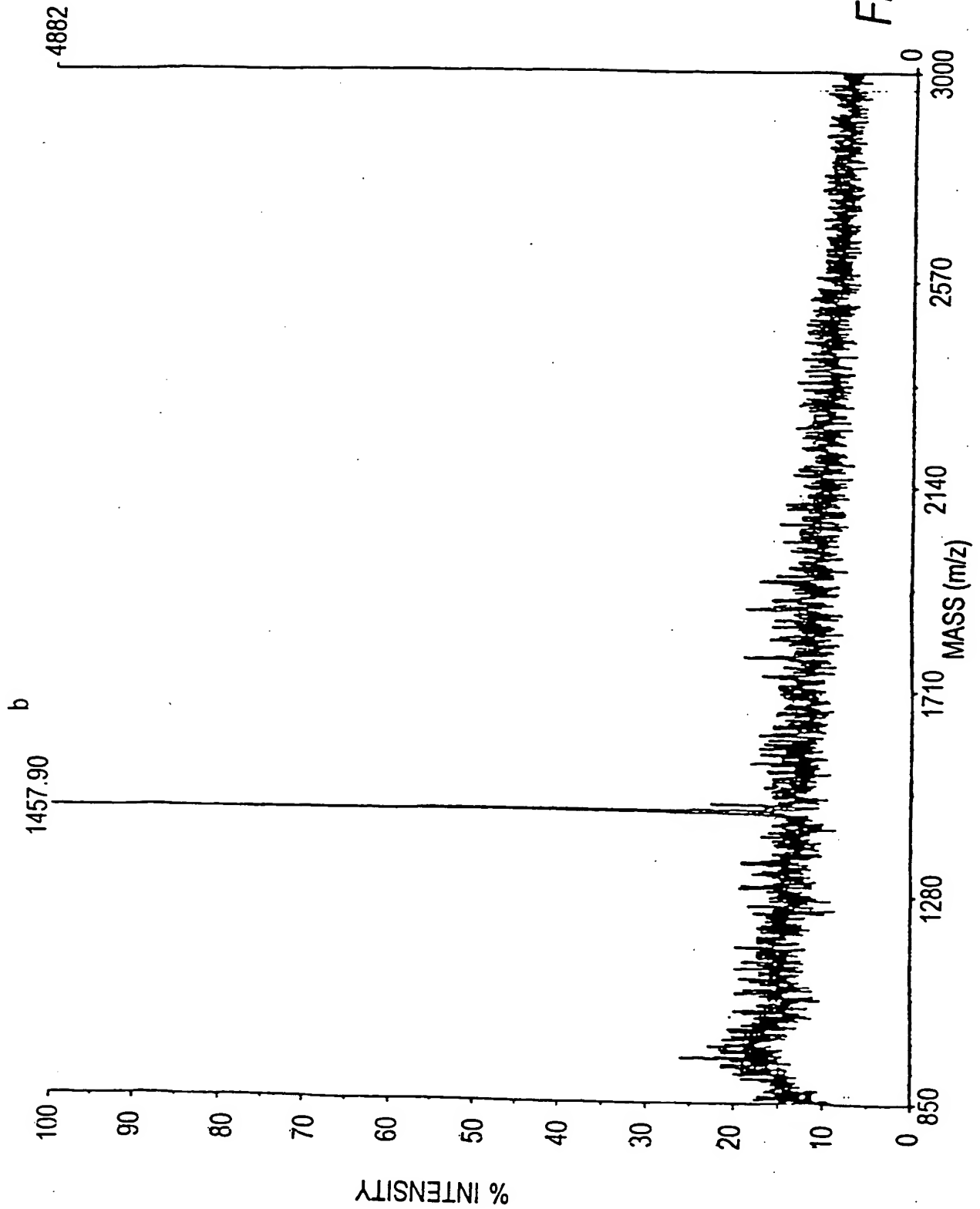


FIG. 10B

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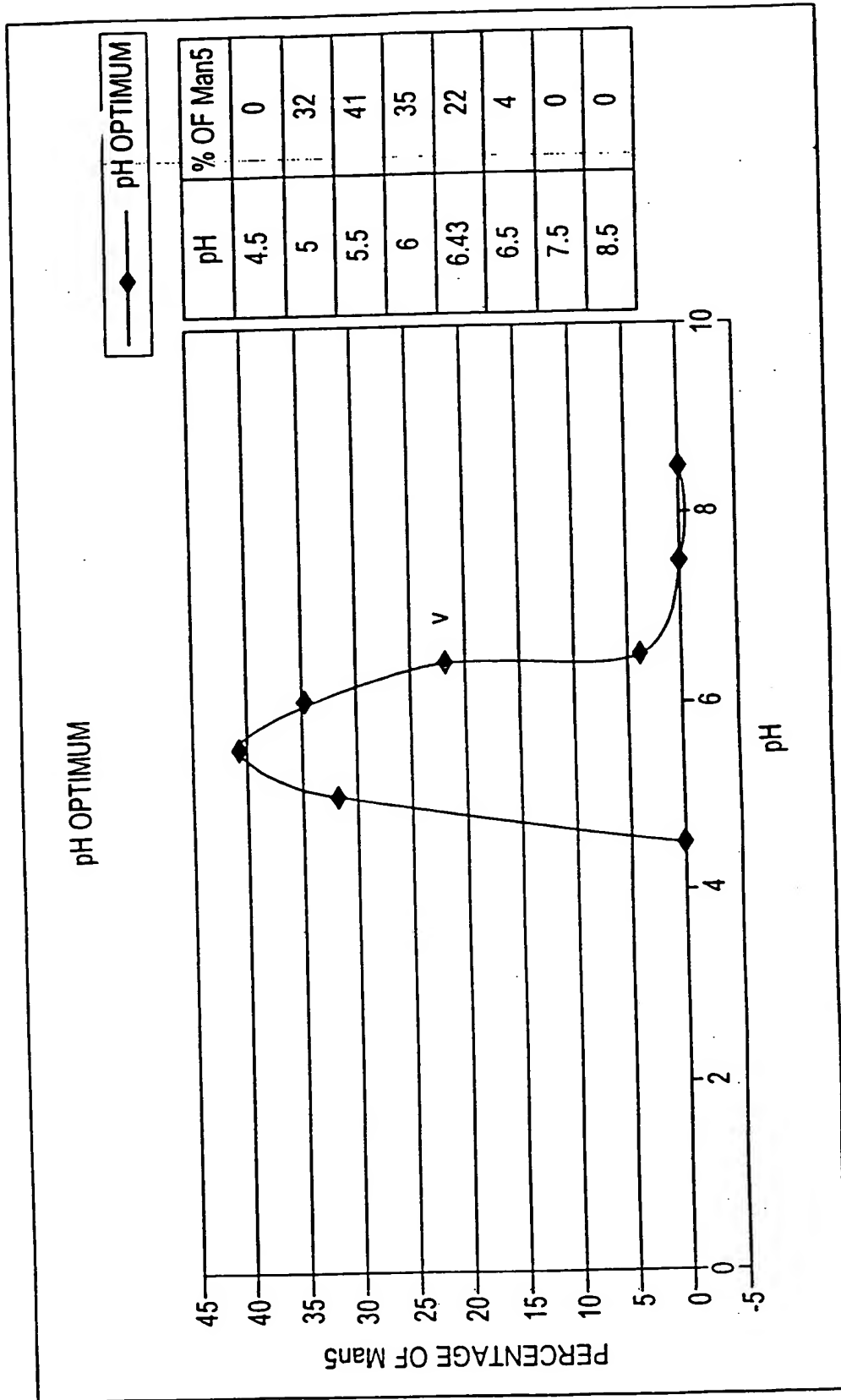
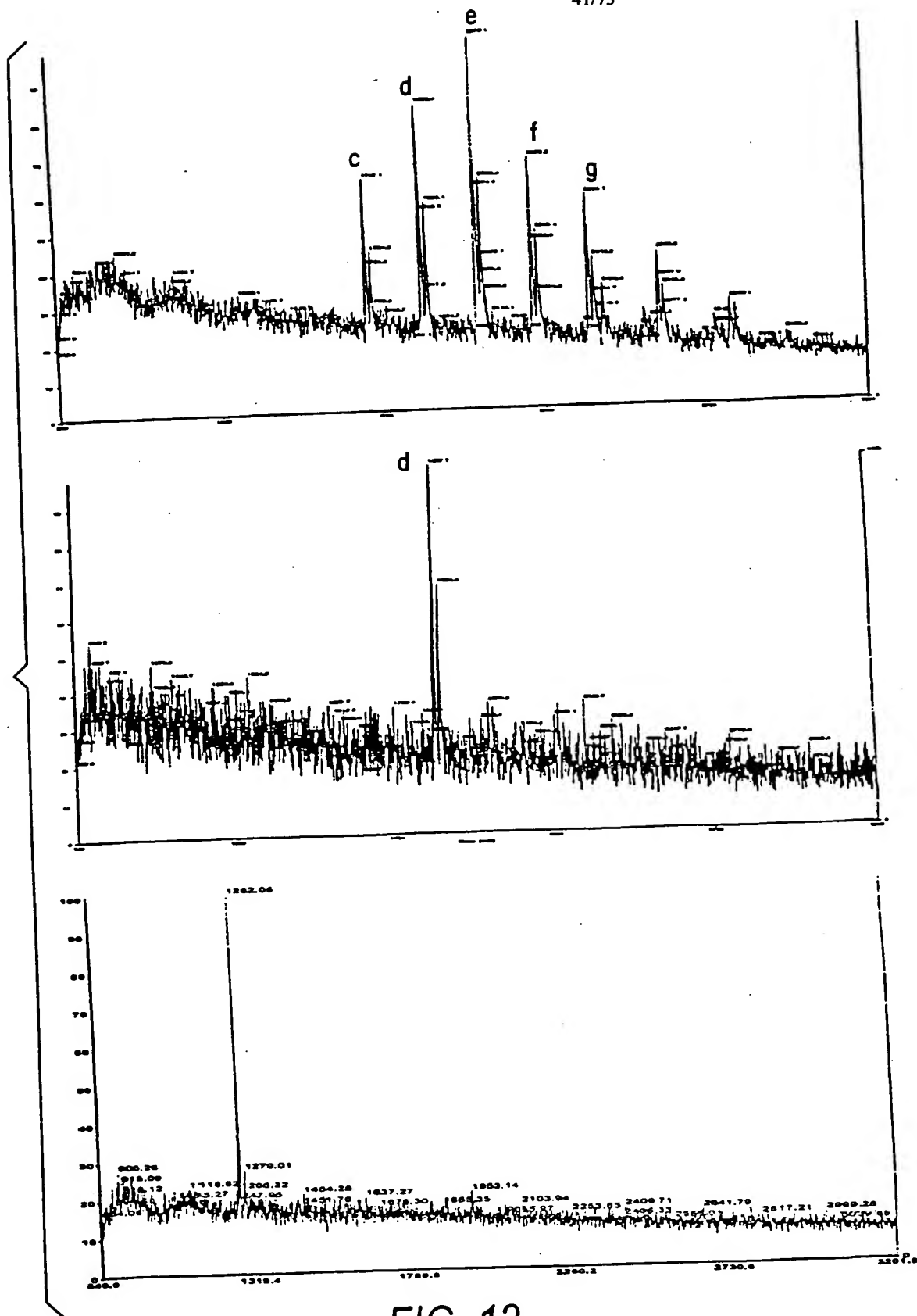


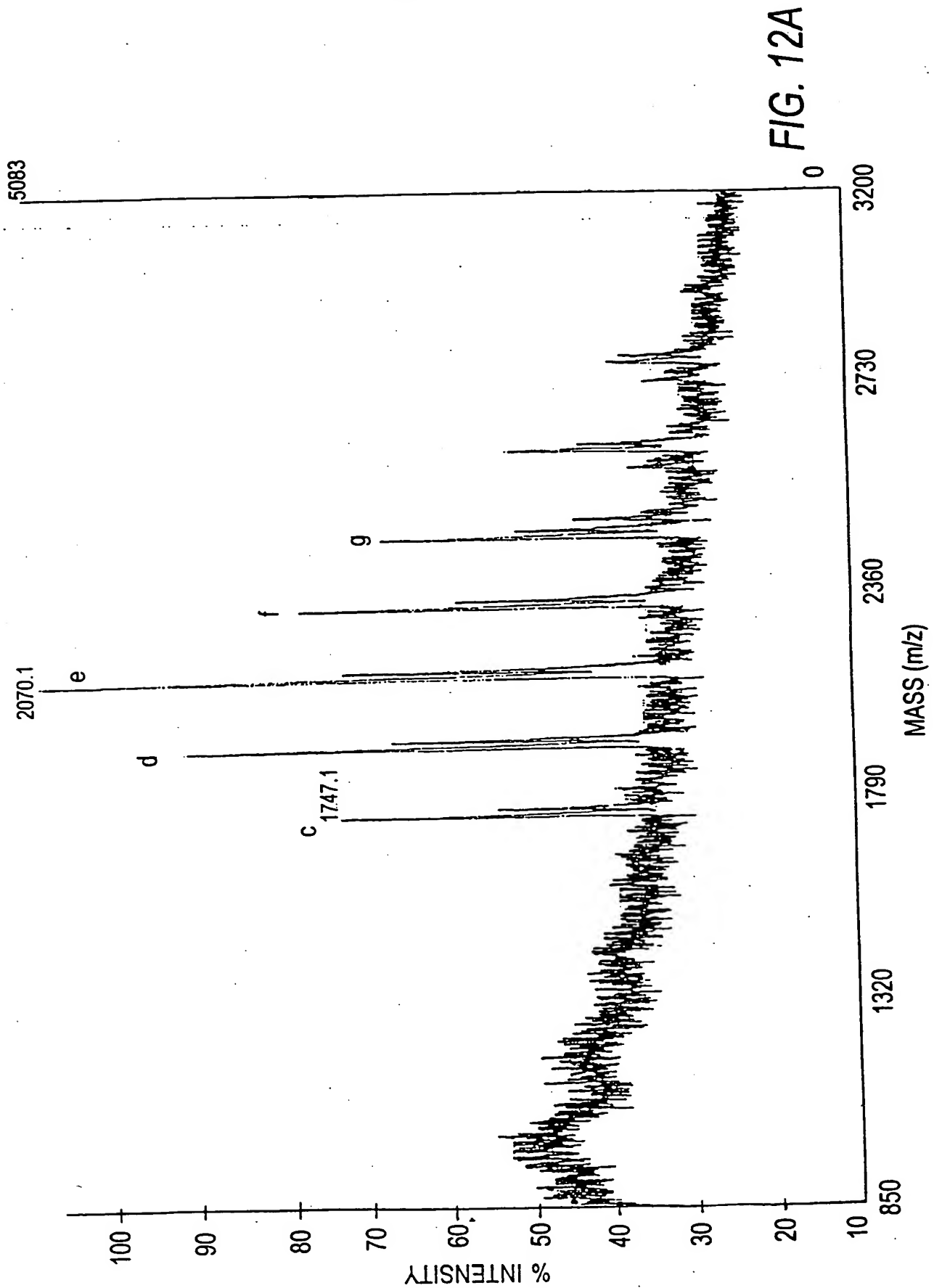
FIG. 11



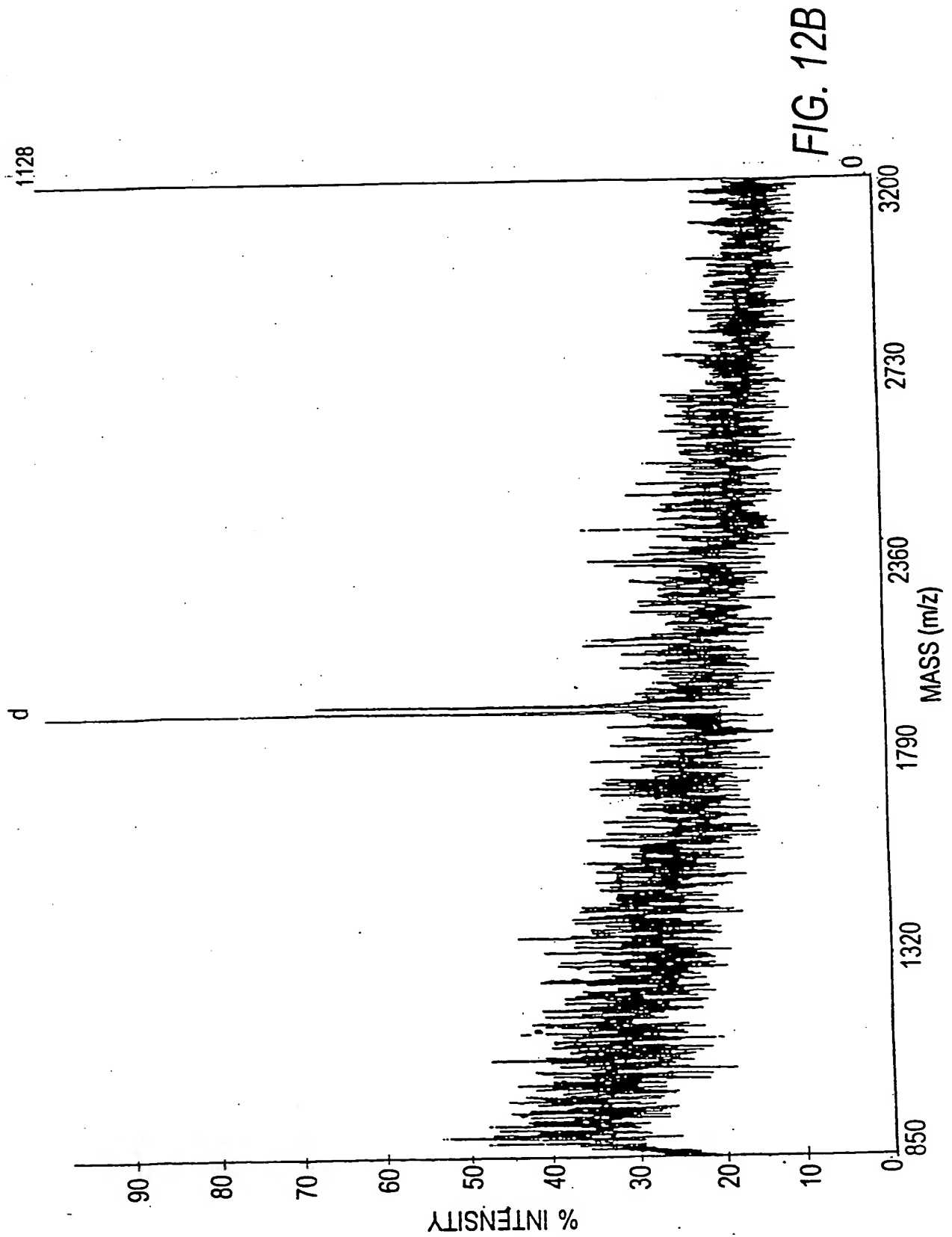
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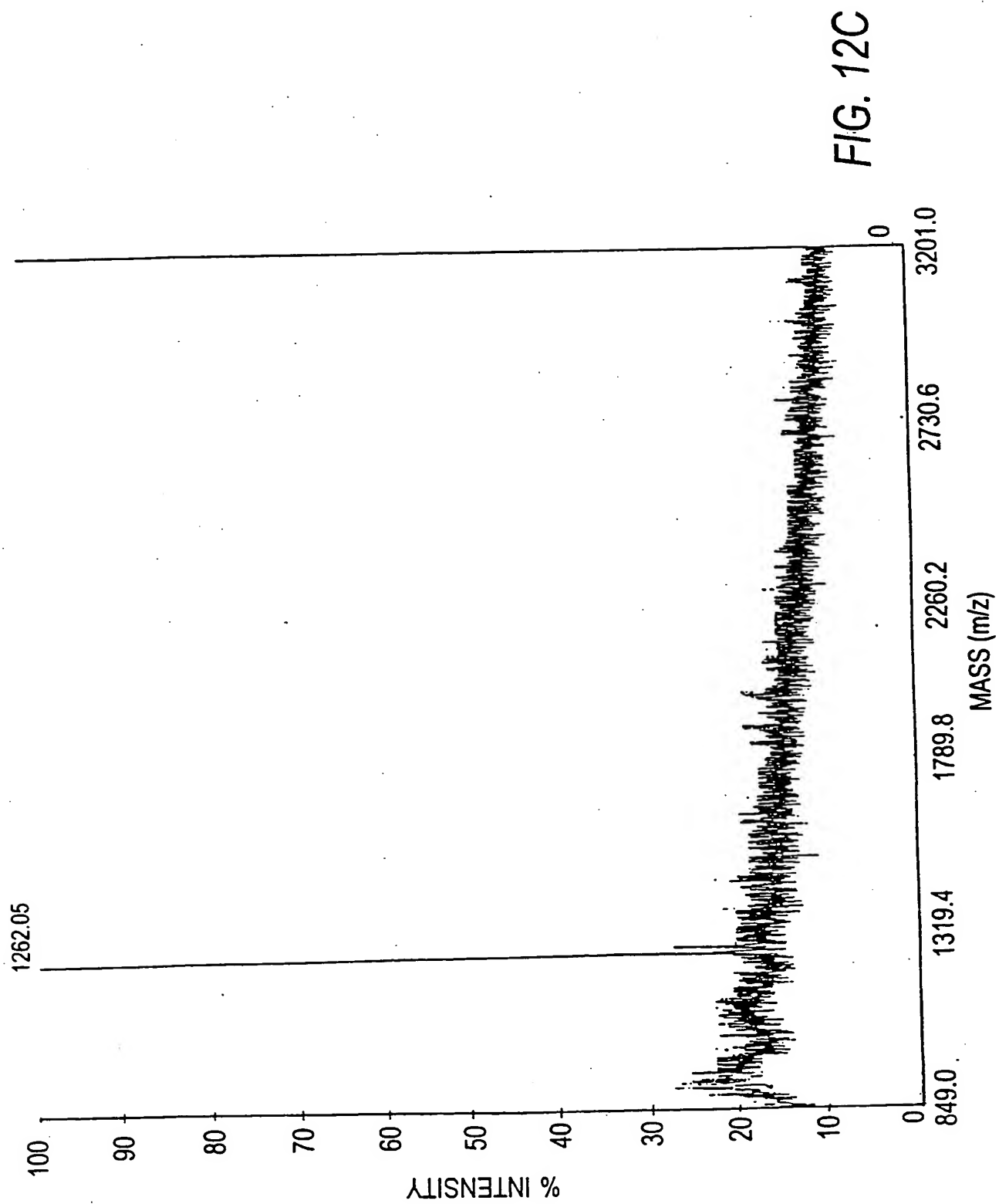
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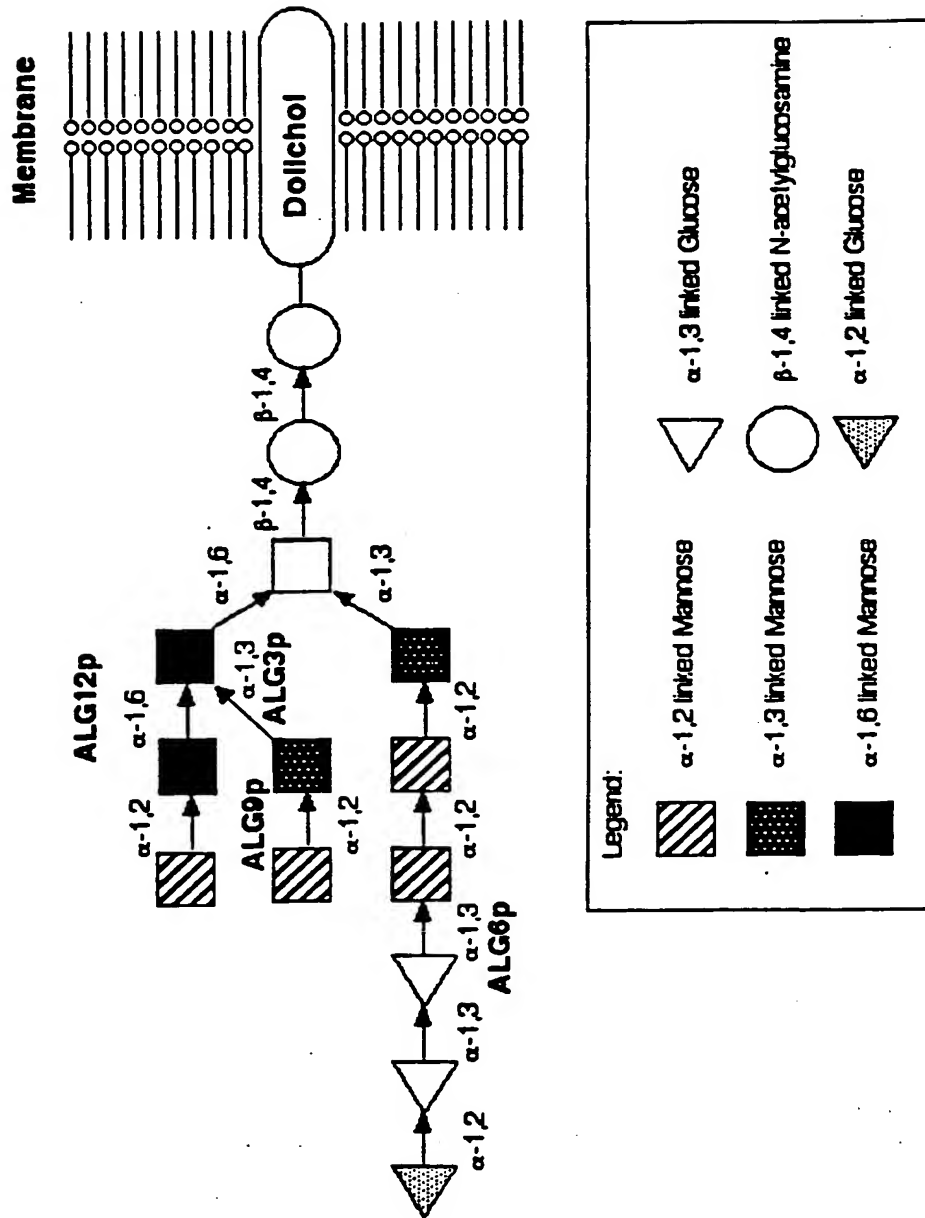


FIG. 13

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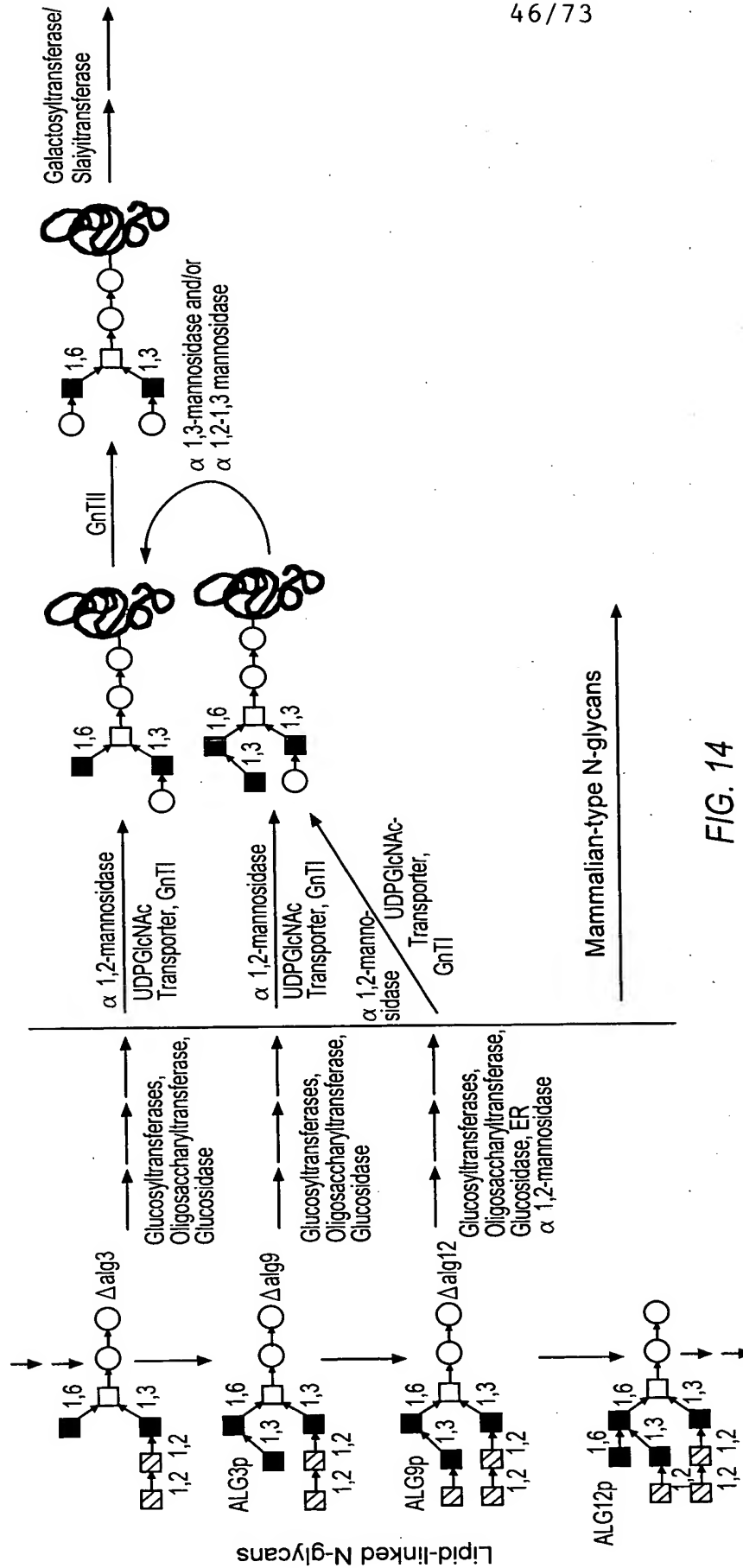


FIG. 14

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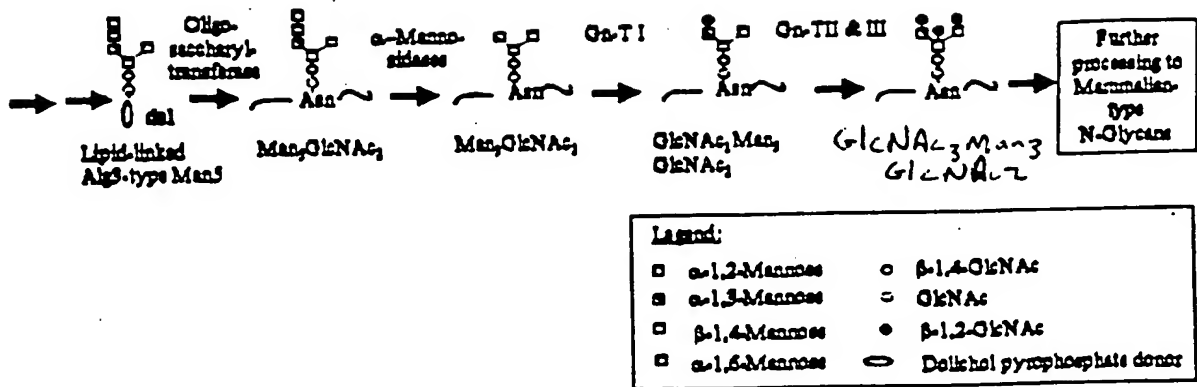


FIG. 15

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# ALG3 Blast 05-22-01

Sequences producing significant alignments:		(bits)	Value
gi 586444 sp P38179 ALG3_YEAST	DOLICHYL-P-MAN:MAN(5)GLCNAC(...797	0.0	
gi 3024226 sp Q92685 ALG3_HUMAN	DOLICHYL-P-MAN:MAN(5)GLCNAC...173	7e-43	
gi 3024221 sp Q24332 NT56_DROVI	LETHAL(2)NEIGHBOUR OF TID P...145	3e-34	
gi 3024222 sp Q27333 NT56_DROME	LETHAL(2)NEIGHBOUR OF TID P...121	3e-27	
gi 10720153 sp P82149 NT53_DROME	LETHAL(2)NEIGHBOUR OF TID ...121	5e-27	
gi 1707982 sp P40989 GLS2_YEAST	1,3-BETA-GLUCAN SYNTHASE CO... 32	2.8	
gi 1346146 sp P38631 GLS1_YEAST	1,3-BETA-GLUCAN SYNTHASE CO... 31	6.6	

## Alignments

### Yeast

>gi|586444|sp|P38179|ALG3\_YEAST DOLICHYL-P-MAN:MAN(5)GLCNAC(2)-PP-DOLICHYL MANNOSYLTRANSFERASE  
(DOL-P-MAN DEPENDENT ALPHA(1-3)-MANNOSYLTRANSFERASE)  
(HM-1 KILLER TOXIN RESISTANCE PROTEIN)  
Length = 458

Score = 797 bits (2059), Expect = 0.0  
Identities = 422/458 (92%), Positives = 422/458 (92%)

```

Query: 1  MEGEQSPQGEKSLQKQKQFVRPPLDLWQDLKDGVRVYVIFDCRANLIVMPLLILFESMLCKI 60
          MEGEQSPQGEKSLQKQKQFVRPPLDLWQDLKDGVRVYVIFDCRANLIVMPLLILFESMLCKI
Sbjct: 1  MEGEQSPQGEKSLQKQKQFVRPPLDLWQDLKDGVRVYVIFDCRANLIVMPLLILFESMLCKI 60

Query: 61  IIKKVAYTEIDYKAYMEQIEMIQLDGMLDYSQVSGGTGPLVYPAGHVLIYKMMYWLTEGM 120
          IIKKVAYTEIDYKAYMEQIEMIQLDGMLDYSQVSGGTGPLVYPAGHVLIYKMMYWLTEGM
Sbjct: 61  IIKKVAYTEIDYKAYMEQIEMIQLDGMLDYSQVSGGTGPLVYPAGHVLIYKMMYWLTEGM 120

Query: 121 DHVERGQVFFRYLYLLTLALQMACYLLHLPPWCVVLAACLSKRLHSIYVLRFLNDCFTTL 180
          DHVERGQVFFRYLYLLTLALQMACYLLHLPPWCVVLAACLSKRLHSIYVLRFLNDCFTTL
Sbjct: 121 DHVERGQVFFRYLYLLTLALQMACYLLHLPPWCVVLAACLSKRLHSIYVLRFLNDCFTTL 180

Query: 181 FMVVTVLGAIVASRCHQRPKLKSLALVISATYSMAVSIKMNALLYFPAMMISLFILNDA 240
          FMVVTVLGAIVASRCHQRPKLKSLALVISATYSMAVSIKMNALLYFPAMMISLFILNDA
Sbjct: 181 FMVVTVLGAIVASRCHQRPKLKSLALVISATYSMAVSIKMNALLYFPAMMISLFILNDA 240

Query: 241 NVILTLLDLVAMIAWQVAVAVPFLRSFPQQYLHCAFNFGRKFMYQWSINWQMMDEEAFND 300
          NVILTLLDLVAMIAWQVAVAVPFLRSFPQQYLHCAFNFGRKFMYQWSINWQMMDEEAFND
Sbjct: 241 NVILTLLDLVAMIAWQVAVAVPFLRSFPQQYLHCAFNFGRKFMYQWSINWQMMDEEAFND 300

Query: 301 KRFXXXXXXXXXXXXXXXXXXSVTRYPRILPDLWSSLCHPLRKNVLANANPAKTIPFVLIASN 360
          KRF                      SVTRYPRILPDLWSSLCHPLRKNVLANANPAKTIPFVLIASN
Sbjct: 301 KRFXHLALLISHLIALTTLFVTRYPRILPDLWSSLCHPLRKNVLANANPAKTIPFVLIASN 360

Query: 361 FIGVLFSSRLHYQFLSWYHWTLPILIFWSGMPFFVGPIWYVLHEWCWNSYPPNSQXXXXX 420
          FIGVLFSSRLHYQFLSWYHWTLPILIFWSGMPFFVGPIWYVLHEWCWNSYPPNSQ
Sbjct: 361 FIGVLFSSRLHYQFLSWYHWTLPILIFWSGMPFFVGPIWYVLHEWCWNSYPPNSQASTLL 420

Query: 421 XXXXXXXXXXXXXXXXXXXXSGSVALAKSHLRTTSSMEKKLN 458
          SGSVALAKSHLRTTSSMEKKLN
Sbjct: 421 LALNTVLLLLLALTLQSGSVALAKSHLRTTSSMEKKLN 458

```

FIG. 16-1



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## Human

>gi|3024226|sp|Q92685|ALG3\_HUMAN DOLICHYL-P-MAN:MAN(5)GLCNAC(2)-PP-DOLICHYL  
MANNOSYLTRANSFERASE

(DOL-P-MAN DEPENDENT ALPHA(1-3)-MANNOSYLTRANSFERASE)  
(NOT56-LIKE PROTEIN)

Length = 438

Score = 173 bits (439), Expect = 7e-43  
Identities = 133/396 (33%), Positives = 195/396 (48%), Gaps = 28/396 (7%)

Query: 26 WQDLKDGVRVIFDCRANLIVMPLLLILFESMLCKIIKKVAYTEIDYKAYMEQIEMIQLD 85  
WQ+ R ++ + R L+V L L E + +I +VAYTEID+KAYM ++E + ++  
Sbjct: 29 WQER----RLLLREPRYTLLVAACLCLEAVGITFWIHRVAYTEIDWKAYMAEVEGV-IN 83

Query: 86 GMLDYSQVSGGTGPLVYPAGHVLIYKMMYWLTEGMDHVERGQVFFRYLYLLTLALQMACY 145  
G DY+Q+ G TGPLVYPAG V I+ +Y+ T + Q F LYL TL L Y  
Sbjct: 84 GTYDYTLQOGDTGPLVYPAGFVYIFMGLYYATSRGTDIRMAQNIFAVLYLATLLLVFLIY 143

Query: 146 Y-LLHLPWC-VVLACLSKRLHSIYVLRLEFNDCTTFLMVVTVLGAIVASRCHQRPKLKK 203  
+ +PP+ + C S R+HSI+VLRLEFND + + +L + QR  
Sbjct: 144 HQTCCKVPFVFFFMCCASYRVHSIFVLRLEFNDP-----VAMVLLFLSINLLLAQRWGWG- 197

Query: 204 SLALVISATYSMAVSIKMNALLYFPAMMISLFILNDANVILTLLDLVAMIAWQVAVAVPF 263  
+S+AVS+KMN LL+ P ++ L L L + A + QV + +PF  
Sbjct: 198 -----CCFFSLAVSVKMNVLFLFAPGLLFLLLTQFGFRGALPKLGICAGL--QVVLGLPF 249

Query: 264 LRSFPQQYLHCAFNFRKFMYSINWQMMDEEAFNDKRFXXXXXXXXXXXXXXXXXVFTRY 323  
L P YL +F+ GR+F++ W++NW+ + E F + F + R+  
Sbjct: 250 LLENPSGYLSRSFDLGRQFLHWTNVNRFLEALFLHRAFLALLTAHLTLLLLFALCRW 309

Query: 324 PRILPDLWSSLCHPLRKNVNLNANPAKTIPFVLIASNFIGVLFSSRLHYQFLSWYHWTLP 383  
R + S L P ++ I L SNFIG+ FSRSLHYQF WY TLP  
Sbjct: 310 HRTGESILSLRDPKRVPPQPLTPNQIVSTLFTSNFIGICFSRSLHYQFVWYFHTLP 369

Query: 384 ILIF-----WSGMPFFVGPIWYVLHEWCWNSYPPNS 414  
L++ W + + + E WN+YP S  
Sbjct: 370 YLLWAMPARWLTHLLRLLVLGLI--ELSWNTYPSTS 403

## Drosophila Vi

>gi|3024221|sp|Q24332|NT56\_DROVI LETHAL(2)NEIGHBOUR OF TID PROTEIN (NOT58)  
Length = 526

Score = 145 bits (366), Expect = 3e-34  
Identities = 103/273 (37%), Positives = 157/273 (56%), Gaps = 17/273 (6%)

Query: 33 VRVYIFDCRANLIVMPLLLILFESMLCKIIKKVAYTEIDYKAYMEQIEMIQLDGMGLDYSQ 92  
++Y+ F+ A IV L++L E+++ ++I++V YTEID+KAYM++ E L+G +YS  
Sbjct: 34 IKYLAFEPAALPIVSVLIVLAEAVINVLVIQVRVPEYTEIDWKAYMQECEGF-LNGTTNYSL 92

Query: 93 VSGGTGPLVYPAGHVLIYKMMYWLTEGMDHVERGQVFFRYLYLLTLALQMACYLLH-LP 151  
+ G TGPLVYPA V IY +Y+LT +V Q F +YLL + L + Y +P  
Sbjct: 93 LRGDTGPLVYPAAFVYIYSGLYLTGQGTNVRLAQYIFACIYLLQMCLVLRLYTKSRKVP 152

Query: 152 PWCVVLACL-SKRLHSIYVLRLEFNDCTTFLMVVTVLGAIVASRCHQRPKLKKSALVIS 210  
P+ +VL+ S R+HSIYVLRLEFND L +L A + QR L S  
Sbjct: 153 PYVLVLSAFTSYRIHSIYVLRLEFNDPVAL-----LLYAALNLFQDQRTLG-----S 200

Query: 211 ATYSMAVSIKMNALLYFPAMMISLFILNDANVILTLLDLVAMIAWQVAVAVPFLRSFPQQ 270  
YS+AV +KMN + A + LF L + V+ TL+ L Q+ + PFLR+ P +  
Sbjct: 201 ICYSLAVGVKMN--ILLFAPALLLFYLANLGVRLTLVQLTICAVLQFLGAPFLRTHPME 258

FIG. 16-2

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Query: 271 YLHCAFNFGRKFMYQWSINWQMMDEEAFNDKRF 303  
YL +F+ GR F ++W++N++ + +E F + F  
Sbjct: 259 YLRGSFDLGRIFEHKWTVNYRFLSKELFEQREF 291

Score = 53.3 bits (127), Expect = 1e-06  
Identities = 31/62 (50%), Positives = 41/62 (66%), Gaps = 6/62 (9%)

Query: 352 IPFVLIASNFIGVLFPSRSLHYQFLSWYHWTLPILIFWSGMPFFVGPIWYVLH--EWCWNS 409  
+PF L NFIGV +RSLHYQF WY +LP L+ WS P+ +G + +L E+CWN+  
Sbjct: 412 LPFFL--CNFIGVACARSLSHYQFYIWFHSLPYLV-WS-TPYSLGVRYLILGIEYCWN 467

Query: 410 YP 411  
YP

Sbjct: 468 YP 469

# *Drosophila melanogaster*

>gi|3024222|sp|Q27333|NT56\_DROME LETHAL(2)NEIGHBOUR OF TID PROTEIN (NOT56)  
(NOT45)

Length = 510

Score = 121 bits (305), Expect = 3e-27  
Identities = 96/272 (35%), Positives = 154/272 (56%), Gaps = 17/272 (6%)

Query: 34 RYVIFDCRANLIVMPLILFESMLCKIIKKVAYTEIDYKAYMEQIEMIQLDGMLDYSQV 93  
+Y++ + A IV ++L E ++ ++I++V YTEID+ AYM++ E L+G +YS +  
Sbjct: 36 KYLLLEPAALPIVGLFVLLAELVINVVVVIQRPVYTEIDWVAYMQECEGF-LNGTTNYSLL 94

Query: 94 SGGTGPLVYPAGHVLIYQMYWLTEGMDHVERGQVFFRYLYLLTLALOMACYLLH-LPP 152  
G TGPLVYPA V IY +Y++T +V Q F +YLL LAL + Y +PP  
Sbjct: 95 RGDGTGPLVYPAAFVYIYSALYYVTSHGTINVRLAQYIFAGIYLLQLALVLRLYSKSRKVPP 154

Query: 153 WCVVLACL-SKRLHSIYVLRFLFNDCTTILFMVTVLGAIVASRCHQRPKLKKSALVISA 211  
+ +VL+ S R+HSIYVLRFLFND + V +L A + +R L S  
Sbjct: 155 YVLVLSAFTSYRIHSIYVLRFLFNDP-----VAVLLLYAALNLFLLDRRTLG-----ST 202

Query: 212 TYSMAVSIKMNALLYFPAMMISLFILNDANVILTLLDLVAMIAWQVAVAVPFLRSFPQOY 271  
+S+AV +KMN + A + LF L + ++ T+L L Q+ + PFL + P +Y  
Sbjct: 203 FFSLAVGVKMN--ILLFAPALLLFYLANLGLLRTILQLAVCGVIQLLLGAPFLLLTHPVEY 260

Query: 272 LHCAFNFGRKFMYQWSINWQMMDEEAFNDKRF 303  
L +F+ GR F ++W++N++ + + F ++ F  
Sbjct: 261 LRGSFDLGRIFEHKWTVNYRFLSRDVFENRTF 292

Score = 49.4 bits (117), Expect = 2e-05  
Identities = 27/60 (45%), Positives = 35/60 (58%), Gaps = 2/60 (3%)

Query: 352 IPFVLIASNFIGVLFPSRSLHYQFLSWYHWTLPILIFWSGMPFFVGPIWYVLHEWCWNSYP 411  
+PF L N +GV SRSLHYQF WY +LP L + + V + L E+CWN+YP  
Sbjct: 407 LPFFL--CNLVGVACSRSLHYQFYVWYFHSPLAWSTPYSLGVRCILGLIEYCWN 464

FIG. 16-3

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Matrix: BLOSUM62  
Gap Penalties: Existence: 11, Extension: 1  
Number of Hits to DB: 28883317  
Number of Sequences: 96469  
Number of extensions: 1107545  
Number of successful extensions: 2870  
Number of sequences better than 10.0: 16  
Number of HSP's better than 10.0 without gapping: 5  
Number of HSP's successfully gapped in prelim test: 11  
Number of HSP's that attempted gapping in prelim test: 2839  
Number of HSP's gapped (non-prelim): 23

length of query: 458  
length of database: 35,174,128  
effective HSP length: 45  
effective length of query: 413  
effective length of database: 30,833,023  
effective search space: 12734038499  
effective search space used: 12734038499  
T: 11  
A: 40  
X1: 15 ( 7.1 bits)  
X2: 38 (14.6 bits)  
X3: 64 (24.7 bits)  
S1: 40 (21.8 bits)  
S2: 67 (30.4 bits)

FIG.16-4

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*S. cerevisiae* ALG3

ATGGAAGGTGAACAGTCTCCGCAAGGTGAAAAGTCTCTGCAAAGGAAGC  
AATTTGTCAGACCTCCGCTGGATCTGTGGCAGGATCTCAAGGACGGTGTG  
CGCTACGTGATCTTCGATTGTAGGGCCAATCTTATCGTTATGCCCCTTTTG  
ATTTTGTTGAAAGCATGCTGTGCAAGATTATCATTAAAGAAGGTAGCTTAC  
ACAGAGATCGATTACAAGGCGTACATGGAGCAGATCGAGATGATTCAGCT  
CGATGGCATGCTGGACTACTCTCAGGTGAGTGGTGGAAACGGGCCCGCTGG  
TGTATCCAGCAGGCCACGTCTTGATCTACAAGATGATGTACTGGCTAACA  
GAGGGAATGGACCACGTTGAGCGCGGGCAAGTGTTTTTCAGATACTTGTA  
TCTCCTTAACTGGCGTTACAAATGGCGTGTTACTACCTTTTACATCTACC  
ACCGTGGTGTGTGGTCTTGGCGTGCCTCTCTAAAAGATTGCACTCTATTTA  
CGTGCTACGGTTATTCAATGATTGCTTCACTACTTTGTTTATGGTCGTCACG  
GTTTTGGGGGCTATCGTGGCCAGCAGGTGCCATCAGCGCCCCAAATTAAA  
GAAGTCCCTTGCGCTGGTGTATCTCCGCAACATACAGTATGGCTGTGAGCA  
TTAAGATGAATGCGCTGTTGTATTTCCCTGCAATGATGATTTCTCTATTCAT  
CCTTAATGACGCGAACGTAATCCTTACTTTGTTGGATCTCGTTGCGATGAT  
TGCAATGGCAAGTCGCAGTTGCAGTGCCCTTCCCTGCGCAGCTTTCCGCAACA  
GTACCTGCATTGCGCTTTTAATTTCCGCGAGGAAGTTTATGTACCAATGGAG  
TATCAATTGGCAAATGATGGATGAAGAGGCTTTCAATGATAAGAGGTTCC  
ACTTGCCCTTTTAATCAGCCACCTGATAGCGCTCACCACACTGTTCTGTC  
CAAGATACCCTCGCATCCTGCCCCGATTTATGGTCTTCCCTGTGCCATCCGC  
TGAGGAAAAATGCAGTGCTCAATGCCAATCCCGCCAAGACTATTCCATTC  
GTTCTAATCGCATCCAACCTTCATCGGCGTCCTATTTTCAAGGTCCCTCCAC  
TACCAGTTTCTATCCTGGTATCACTGGACTTTGCCTATACTGATCTTTTGGT  
CGGGAATGCCCTTCTTCGTTGGTCCCATTGGTACGTCTTGACAGAGTGGT  
GCTGGAATTCTATCCACCAAACCTCACAAGCAAGCACGCTATTGTTGGCA  
TTGAATACTGTTCTGTTGCTTCTATTGGCCTTGACGCAGCTATCTGGTTCGG  
TCGCCCTCGCCAAAAGCCATCTTCGTACCACCAGCTCTATGGAAAAAAG  
CTCAACTGA

*S. cerevisiae* Alg3p

MEGEQSPQGEKSLQRKQFVRPPLDLWQDLKDGVRVYVIFDCRANLIVMPLLIL  
FESMLCKIIKKVAYTEIDYKAYMEQIEMIQLDGM LDYSQVSGGTGPLVYPAG  
HVLIYKMMYWLTEGMDHVERGQVFFRYLYLLTLALQMACYLLHLPPWCV  
VLACLSKRLHSIYVLRLFNDCFTTLFMVVTVLGAIVASRCHQRPKLKKSALV  
ISATYSMAVSIMNALLYFPAMMISLFILNDANVILTLLDLVAMIAWQVAVA  
VPFLRSFPQQYLHCAFNFGRKFMYQWSINWQMMDEEAFNDKRFHLALLISHL  
IALTTLFVTRYPRILPDLWSSLCHPLRKNVNLNANPAKTIPFVLIASNFIGVLFS  
RSLHYQFLSWYHWTLPIILFWSGMPFFVGPIWYVLHEWCWNSYPPNSQASTL  
LLALNTVLLLLLALTQLSGSVALAKSHLRTTSSMEKKLN

FIG. 17

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*P. pastoris* ALG3

ATGCCTCCGATAGAGCCAGCTGAAAGGCCAAAGCTTACGCTGAAAAATGT  
TATCGGTGATCTAGTGGCTCTTATTCAAACGTTTTATTTAACCCAGATTTT  
AGTGTCTTCGTTGCACCTCTTTTATGGTTAGCTGATTCCATTGTTATCAAGG  
TGATCATTGGCACTGTTTCTACACAGATATTGATTTTTCTTCATATATGCA  
ACAAATCTTTAAAATTTCGACAAGGAGAATTAGATTATAGCAACATATTTG  
GTGACACCGGTCCATTGGTTTACCCAGCCGGCCATGTTTCATGCTTACTCAG  
TACTTTCGTGGTACAGTGATGGTGGAGAAGACGTCAGTTTCGTTCAACAA  
GCATTTGGTTGGTTATACCTAGGTTGCTTGTTACTATCCATCAGCTCCTACT  
TTTTCTCTGGCTTAGGGAAAATACCTCCGGTTTATTTTGTGTTTGTGGTAGC  
GTCCAAGAGACTGCATTCAATATTTGTATTGAGACTCTTCAATGACTGTTT  
AACAACATTTTTGATGTTGGCAACTATAATCATCCTTCAACAAGCAAGTAG  
CTGGAGGAAAGATGGCACAACCTATTCCATTATCTGTCCCTGATGCTGCAG  
ATACGTACAGTTTAGCCATCTCTGTAAAGATGAATGCGCTGCTATACCTCC  
CAGCATTCCACTACTCATATATCTCATTGTGACGAAAATTTGATTAAAG  
CCTTGGCACCTGTTCTAGTTTTGATATTGGTGCAGTAGGAGTCGGTTATT  
CGTTCATTTTACCGTTGCACTATGATGATCAGGCAAATGAAATTCGTTCTG  
CCTACTTTAGACAGGCTTTTGACTTTAGTCGCCAATTTCTTTATAAGTGGA  
CGGTTAATTGGCGCTTTTTGAGCCAAGAACTTTCAACAATGTCCATTTTC  
ACCAGCTCCTGTTTGCTCTCCATATTATTACGTTAGTCTTGTTTCATCCTCAA  
GTTCTCTCTCCTAAAAACATTGGAAAACCGCTTGGTAGATTGTGTTGGA  
CATTTTCAAATTTTGGAAGCCAACCTTATCTCCAACCAATATTATCAACGA  
CCCAGAAAGAAGCCCAGATTTTGTTTACACCGTCATGGCTACTACCAACTT  
AATAGGGGTGCTTTTTGCAAGATCTTTACACTACCAGTTCCTAAGCTGGTA  
TGCGTTCTCTTGCCATATCTCCTTTACAAGGCTCGTCTGAACCTTATAGCA  
TCTATTATTGTTTATGCCGCTCACGAGTATTGCTGGTTGGTTTTCCAGCTA  
CAGAACAAAGTTCGCGCTTGTGGTATCTATCTTACTACTTATCCTGATTC  
TCATTTTTACCAACGAACAGTTATTTCTTCTCAATCGGTCCCTGCAGAAA  
AAAAGAATACATAA

*P. pastoris* Alg3p

MPPIEPAERP KLT LKNVIGDLVALIQNVLFNPDFS VFVAPLLWLADSI VIKVII G  
TVSYTDIDFSSYMQQIFKIRQGELDYSNIFGDTGPLVYPAGHVHAYSVLSWYS  
DGGEDVSFVQQAFGWLYLGCLLSISSYFFSGLGKIPPVYFVLLVASKRLHSIF  
VLRLFNDCLTTFMLATHILQQASSWRKDGTTPLSVPDAADTYS LAISVKMN  
ALLYLPAFLLLIYLICDENLIKALAPVLVLILVQVGVGYSFILPLHYDDQANEIR  
SAYFRQAFDFSRQFLYKWTNVNWRFLSQETFNNVHFHQLLFALHIITLVLFILKF  
LSPKNIGKPLGRFVLDFKFWKPTLSPTNIINDPERSPDFVYTVMATTNLIGVLF  
ARSLHYQFLSWYAFSLPYLLYKARLNFIASIIVYAAHEYCWLVFPATEQSSAL  
LVSILLILILIFTNEQLFPSQSVP AEKNT

FIG. 18

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*P. pastoris* ALG3 BLAST

Sequences producing significant alignments: (bits) Value

gi 586444 sp P38179 ALG3 YEAST Dolichyl-P-Man:Man(5)GlcNAc(...228	2e-58
gi 12802365 qb AAK07848.1 AF309689.10 putative NOT-56 manno...212	8e-54
gi 984725 qb AAA75352.1  ORF 1 206	4e-52
gi 7492702 pir T39084 probable mannosyltransferase - fissi...176	8e-43
gi 16226531 qb AAL16193.1 AF428424.1 At2g47760/F17A22.15 {A...164	2e-39
gi 25367230 pir B84919 Not56-like protein [imported] - Ara...164	3e-39
gi 25814791 emb CAB70171.2  Hypothetical protein K09E4.2 {C...161	2e-38
gi 17535001 ref NP_496950.1  Putative plasma membrane membr...160	3e-38
gi 1654000 emb CAA70220.1  Not56-like protein [Homo sapiens...155	2e-36
gi 13279206 qb AAH04313.1 AAH04313 Unknown (protein for IMA...154	2e-36
gi 22122365 ref NP_666051.1  hypothetical protein MGC36684 ...150	3e-35
gi 21292031 qb EAA04176.1  agCP3388 [Anopheles gambiae str....120	4e-26
gi 1780792 emb CAA71167.1  lethal(2)neighbour of tid [Droso...114	3e-24

Alignments

*S. cerevisiae*

Score = 228 bits (580), Expect = 2e-58  
Identities = 154/429 (35%), Positives = 229/429 (53%), Gaps = 37/429 (8%)

Query: 9 RPKLT LKNVIGDLVALIQNVLFNPDFSVFVAPLLWLADSIKVIIGTVSYTDIDFSSYM 68  
RP L L DL ++ V+F+ ++ V PLL L +S++ K+II V+YT-ID+ +YM  
Sbjct: 20 RPPLDLWQ---DLKDGVRVYIFDCRANLIVMPLLILFESMLCKIIKKVAYTEIDYKAYM 76

Query: 69 QQIFKIR-QGELDYSNIFGDTGPLVYPAGHVHAYSVLSWYSDGGEDVSFVQQAFGWLYLG 127  
+QI I+ G LDYS + G TGPLVYPAGHV Y ++ W ++G + V Q F +LYL  
Sbjct: 77 EQIEMIQLDGM LDYSQVSGGTGPLVYPAGHVLIYKMYWLTEGMDHVERGQVFFRYLYLL 136

Query: 128 CLLLSISSYFFSGLGKIPPVYFVLLVASKRLHSIFVLRLFNDCLTTFMLLATI---IILQ 184  
L L ++ Y+ L +PP VL SKRLHSI+VLRLFNDC TT M+ T+ I+  
Sbjct: 137 TLALQMACYY---LLHLPWCVVLA LSKRLHSIYVLRLFNDCFTTLMFVVTVLGAIVAS 193

Query: 185 QASSWRKDGTTIPLSVPAADTYS LAISVKMNNXXXXXXXXXXXXXXXXXCDENLIKALAPXX 244  
+ K ++ L + + TYS+A+S+KMN D N+I L  
Sbjct: 194 RCHQRPKLKKS LALVI---SATYSMAVSIKMNALLYFPAMMISLFILNDANVILTLLDLV 250

Query: 245 XXXXXXXXXXXXYSFILPLHYDDQANEIRSAYFRQAFDFSRQFLYKWTNVNRFSLQETFN NV 304  
F+ Y AF+F R+F+Y+W++NW+ + +E FN+  
Sbjct: 251 AMIAWQVAVAVPFL-----RSFPQQLHCAFNFGRKFMYQWSINWQMMDEEAFNDK 301

Query: 305 HFHQLLFALHIITL-VLFILKFLSPKNIGKPLGRFVLDIFKFWKPTLSPTNIIN-DPERS 362  
FH L H+I L LF+ ++ R + D++ L ++N +P ++  
Sbjct: 302 RFHLALLISHLIALTTLFVTRY-----PRILPDLWSSLCHPLRKNAVLNANPAKT 351

Query: 363 PDFVYTVMATTNLIGVLFARSLHYQFLSWYAFSLPYLLYKARLNFIASIIVYAAHEYCW L 422  
F V+ +N IGVLF+RSLHYQFLSWY ++LP L++ + + F I Y HE+CW  
Sbjct: 352 IPF---VLIASNFIGVLFARSLHYQFLSWYHWTLPILIFWSGMPFFVGPIWYVLHEWCWN 408

Query: 423 VFPATEQSS 431  
+P Q+S  
Sbjct: 409 SYPPNSQAS 417

FIG. 19-1

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*Neurospora crassa*

Score = 212 bits (540), Expect = 8e-54  
Identities = 140/400 (35%), Positives = 212/400 (53%), Gaps = 29/400 (7%)

Query: 35 SVFVAPLLWLADSIKVIIGTVSYTDIDFSSYMQQIFKIROGELDYSNIFGDTGPLVYP 94  
S + P L+L D+++ +II V YT+ID+++YM+Q+ +I GE DY+ + G TGPLVYP  
Sbjct: 33 SKLIPPALFLVDALLCGLIIWKVPYTEIDWAAAYMEQVSQILSGERDYTEKVRGGTGPLVYP 92

Query: 95 AGHVHAYSVLSWYSDGGEDVSFVQQAAGWLYLGCLLLSISSYFFSGLGKIPPVYFVLLVA 154  
A HV+ Y+ L +D G ++ QQ F LY+ L + + Y+ K PP F LL  
Sbjct: 93 AAHVYIYTGLYHLTDEGRNILLAQQLFAGLYMVT LAVVMGCYW---QAKAPPYLFPLLT 149

Query: 155 SKRLHSIFVLRLENDCLTTFMLATIIILQQASSWRKDGTTIPLSVDAADTYSLAISVK 214  
SKRLHSIFVLR FNDC + I Q+ +W+ A Y+L + VK  
Sbjct: 150 SKRLHSIFVLRFCFNDCAFVLFWLAIFFQR-RNWQA-----GALLYTLGLGVK 197

Query: 215 MXXXXXXXXXXXXXXXXXCDENLIKALAPXXXXXXXXXXXXXYSFILPLHYDDQANEIRSAY 274  
M + + L F+ HY + Y  
Sbjct: 198 MTLTLLSLPAVGIVLFLGSG-SFVTTLQLVATMGLVQILIGVPFL--AHYPT-----Y 247

Query: 275 FROAFDFSROFLYKWTNVNWRFLSQETFNNVHFHQLLFALHIITLVLFIL-KFLSPKNIGK 333  
+AF+ SRQF +KWTNVNRF+ +E F + F L ALH++ L +FI +++ P K  
Sbjct: 248 LSRAFELSRQFFKWTNVNRFVGEIEFLSKGFALTLLALHVLVLGIFITTRWIKPAR--K 305

Query: 334 PLGRFVLDFKFWKPTLS-PTNIINDPERSPDFVYTVMATTNLIGVLFARSLHYQFLSWY 392  
L + + + KP L+ P + + +P ++ T + + N +G+LFARSLHYQF ++  
Sbjct: 306 SLVQLISFVLLAGKPLTVPFHRAAARDVTPRYIMTTILSANAVGLLFARSLHYQFYAYV 365

Query: 393 AFSLPYLLYKARLNFIASIIVYAAHEYCWLVPATEQSSA 432  
A+S P+LL++A L+ + +++A HE+ W VFP+T SSA  
Sbjct: 366 AWSTPFLWRAGLHPVLVYLLWAVHEWANNVFPSTPASSA 405

*Schizosaccharomyces pombe*

Score = 176 bits (445), Expect = 8e-43  
Identities = 132/390 (33%), Positives = 194/390 (49%), Gaps = 35/390 (8%)

Query: 42 LWLADSIKVIIGTVSYTDIDFSSYMQQIFKIROGELDYSNIFGDTGPLVYPAGHVHAY 101  
L L + + II V YT+ID+ +YM+Q+ GE DY ++ G TGPLVYP GHV Y  
Sbjct: 30 LLLLEIPFVFAIISKVPYTEIDWIAAYMEQVNSFLLGERDYKSLVGCTGPLVYPGGHVFLY 89

Query: 102 SVLSWYSDGGEDVSFVQQAAGWLYLGCLLLSISSYFFSGLGKIPPVYFVLLVASKRLHSI 161  
++L + +DGG ++ Q F ++Y + +I Y F + + P +VLL+ SKRLHSI  
Sbjct: 90 TLLYYLTDGGTNIVRAQYIFAFVY--ITTAIVGYLFK-IVRAPFYIYVLLILSKRLHSI 146

Query: 162 FVLRLENDCLTTFMLATIIILQQASSWRKDGTTIPLSVDAADTYSLAISVKMXXXXXX 221  
F+LRLFND + L + I+ W + A+ S+A SVKM+  
Sbjct: 147 FILRLENDGFNS-LFSSLFILSSCKKKWVR-----ASILLSVACSVKMSSLLYV 194

Query: 222 XXXXXXXXXXXXCDENLIKALAPXXXXXXXXXXXXXYSFILPLHYDDQANEIRSAYFRQAFDF 281  
L++ L P + + + +Y+ QAFDF  
Sbjct: 195 PAYLV-----LLQILGPKKTMHIFVIIIVQILFSIPF----LAYFWSYWTQAFDF 242

Query: 282 SROFLYKWTNVNWRFLSQETFNNVHFHQLLFALHIITLVLFILKFLSPKNIGKPLGRFVLD 341  
R F YKWTNVNRF+ + F + F + LH+ LV F K + + P  
Sbjct: 243 GRAFDYKWTNVNRFIPRSIFESTSFSTSILFLHVALLVAFTCKHWNKLSRATP----- 295

Query: 342 IFKFWKPTLSPTNIINDPERSPDFVYTVMATTNLIGVLFARSLHYQFLSWYAFSLPYLLY 401  
F L+ + +P+P++T +AT+NLI+L ARSLHYQF +W+A+ PYL Y

FIG. 19-2

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Sbjct: 296 -FAMVNSMLTLKPLPKLQLATPNFIFTALATSNLIGILCARSLHYQFYAWFAWYSPYLCY 354

Query: 402 KARLNFIASIIIVYAAHEYCWLVPATEQSS 431  
+A I ++ EY W VFP+T+ SS

Sbjct: 355 QASFPAPIVIGLWMLQEYAWNVPSTKLSS 384  
*Arabidopsis thaliana*

Score = 164 bits (415), Expect = 2e-39  
Identities = 131/391 (33%), Positives = 194/391 (49%), Gaps = 29/391 (7%)

Query: 42 LWLADSIIVIKVIIGTVSYTDIDFSSYMQQIFKIROGELDYSNIFGDTGPLVYPAGHVHAY 101  
L LAD+I++ +II V YT ID+ +YM Q+ GE DY N+ GDTGPLVYPAG ++ Y

Sbjct: 39 LILADAILVALIIAYVPYTKIDWDAYMSQVSGFLGGERDYGNLKGDTGPLVYPAGFLYVY 98

Query: 102 SVLSWYSDGGEDVSFVQQAQFGLYLGLLLSISSYFFSGLGKIPPVYFVLLVASKRLHSI 161  
S + + G +V Q FG LY+ L + + Y + + +P LL SKR+HSI

Sbjct: 99 SAVQNLTTGG--EVYPAQILFGVLYIVNLGIVLIIYVKTDV--VPWWALSLLCLSKRIHSI 154

Query: 162 FVLRLFNDCLTTFMLLATIIILQQASSWRKDGTTIPLSVDAADTYSLAISVKMNXXXXX 221  
FVLRLFNDC L+ A++ + +RK + + +S A+SVKMN

Sbjct: 155 FVLRLFNDCFAMTLLHASMALFL----YRKWHLGMLV-----FSGAVSVKMNVLVLA 202

Query: 222 XXXXXXXXXXXCDENLIKALAPXXXXXXXXXXXXXXXXXSYFILPLHYDDQANEIRSAYFRQAFDF 281  
N+I ++ F++ +Y AFD

Sbjct: 203 PTLLLLLLKAM--NIIGVVSALAGAALAQILVGLPFLITYPV-----SYIANAFDL 251

Query: 282 SRQFLYKWTNVNRFLSQETFNNVHFHQLLFALHIIITLVLFILKFLSPKNIGKPLGRFVLD 341  
R F++ W+VN++F+ + F + F L H+ LV F + K+ G +G

Sbjct: 252 GRVFIHFWSVNFKFVPERVVFVSKEFAVCLLIAHLFLLVAFA-NYKWCKHEGGIIGFMRSR 310

Query: 342 IFKFWKP-TLSPTNIINDPERSPDFVYTMATTNLIGVLFARSLHYQFLSWYAFSLPYLL 400  
F P +LS +++ + + V T M N IG++FARSLHYQF SWY +SLPYLL

Sbjct: 311 HFFLTLPSSLSFSFSDVSASRIITKEHVVTAMFVGNFIGIVFARSLHYQFYSWYFYSLPYLL 370

Query: 401 YKARLNFIASIIIVYAAHEYCWLVPATEQSS 431  
++ +I++ E CW V+P+T SS

Sbjct: 371 WRTFPPTWLRRLIMFLGIELCWNVPSTPSSS 401

FIG. 19-3



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*K. lactis* ALG3

TTTGTTTACAAGCTGATACCAACGAACATGAATACACCGGCAGGTTTACT  
GAAGATTGGCAAAGCTAACCTTTTACATCCTTTTACCGATGCTGTATTACAG  
TGCGATGAGAGTAAACGCAGAACAAATTGCATACATTTTACTTGTTACCA  
ATTACATTGGAGTACTATTTGCTCGATCATTACACTACCAATTCCTATCTT  
GGTACCATTGGACGTTACCAGTACTATTGAATTGGGCCAATGTTCCGTATC  
CGCTATGTGTGCTATGGTACCTAACACATGAGTGGTGCTGGAACAGCTAT  
CCGCCAAACGCTACTGCATCCACACTGCTACACGCGTGTAACACATACTG  
TTATTGGCTGTATTCTTAAGAGGACCCGCAAACCTCGAAAAGTGGTGATAA  
CGAAACAACACACGAGAAAGCTGAG

*K. lactis* Alg3p

FVYKLIPTNMNTPAGLLKIGKANLLHPFTDAVFSAMRVNAEQIAYILLVTNYI  
GVLFARSLHYQFLSWYHWTLPVLLNWANVPYPLCVLWYLTHEWCWNSYPP  
NATASTLLHACNTYCYWLYSZEDPQTRKVVITKQHTRKL

FIG. 20

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# *K. lactis* ALG3 BLAST

Score E (bits) Value  
Sequences producing significant alignments:

qi 586444 sp P38179 ALG3 YEAST	Dolichyl-P-Man:Man(5)GlcNAc(...)	125	1e-28
qi 984725 qb AAA75352.1	ORF 1	94	4e-19
qi 16226531 qb AAL16193.1 AF428424 1	At2g47760/F17A22.15 [A...]	72	1e-12
qi 25367230 pir B84919	Not56-like protein [imported] - Ara...	72	1e-12
qi 21292031 qb EAA04176.1	agCP3388 [Anopheles gambiae str....]	69	2e-11
qi 20892051 ref XP_148657.1	similar to Lethal(2)neighbour ...	65	2e-10

## Alignments

### *S. cerevisiae*

Score = 125 bits (314), Expect = 1e-28  
Identities = 60/120 (50%), Positives = 83/120 (69%), Gaps = 1/120 (0%)  
Frame = +3

Query: 66 ANLLHPFT-DAVFSAMRVNAEQIAYILLVTNYIGVLFARSLHYQFLSWYHWTLPVLLNWA 242  
++L HP ++AV +A A+ I ++L+ +N+IGVLF+RSLHYQFLSWYHWTLP+L+ W+  
Sbjct: 332 SSLCHPLRKNVAVLNANP--AKTIPFVLIASNFIVGLFSRSLHYQFLSWYHWTLPILIFWS 389

Query: 243 NVYPYPLCVLWYLTHEWCWNSYPPNATASTLLHACNTYCYWLYS\*EDPQTRKVVITKQHTR 422  
+P+ + +WY+ HEWCWNSYPPN+ ASTLL A NT L + + V + K H R  
Sbjct: 390 GMPFFVGPIWYVLHEWCWNSYPPNSQASTLLALNTVLLLLLA-LTQLSGSVALAKSHLR 448

### *A. thaliana*

Score = 72.0 bits (175), Expect = 1e-12  
Identities = 42/107 (39%), Positives = 57/107 (53%), Gaps = 3/107 (2%)  
Frame = +3

Query: 84 FTDVFSAMRVNAEQIAYILLVTNYIGVLFARSLHYQFLSWYHWTLPVLLNWANVYPYPLC 263  
F+D S + + E + + V N+IG++FARSLHYQF SWY ++LP LL P L  
Sbjct: 322 FSDVSASRI-ITKEHVVTAMFVGNFIGIVFARSLHYQFYFYSLPYLLWRTPFPTWLR 380

Query: 264 VLWYLTHEWCWNSYPPNATASTL---LHACNTYCYWLYS\*EDPQTRK 395  
++ +L E CWN YP ++S L LH WL DP K  
Sbjct: 381 LIMFLGIELCWNVYPSTPSSSGLLLCLHLIILVGLWLAPSVDPYQLK 427

FIG. 21

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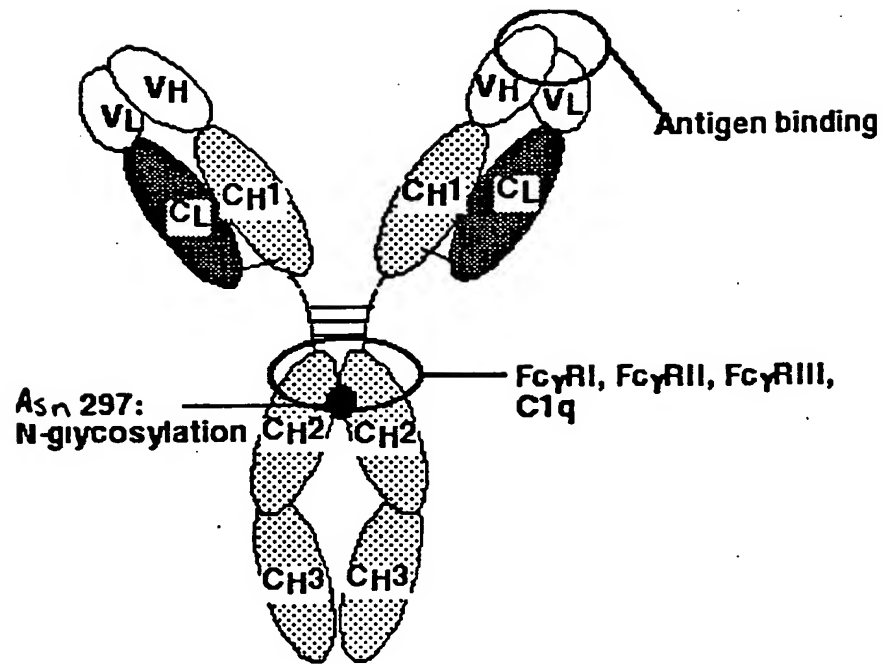


FIG. 22

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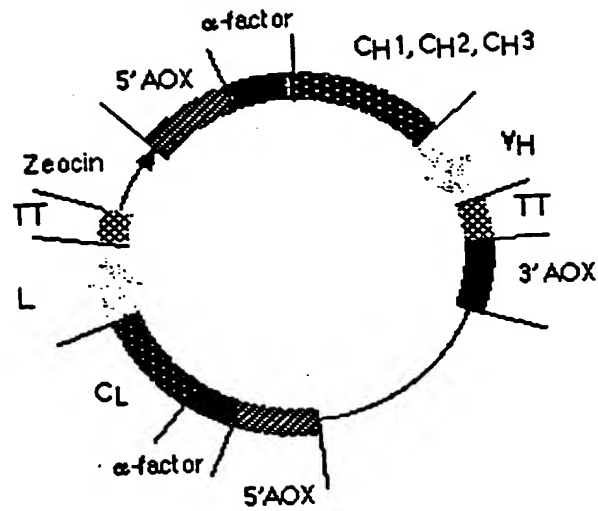


FIG. 23

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>gi|6754685|ref|NM\_010795.1| Mus musculus mannoside acetyl  
glucosaminyltransferase 3 (Mgat3), mRNA

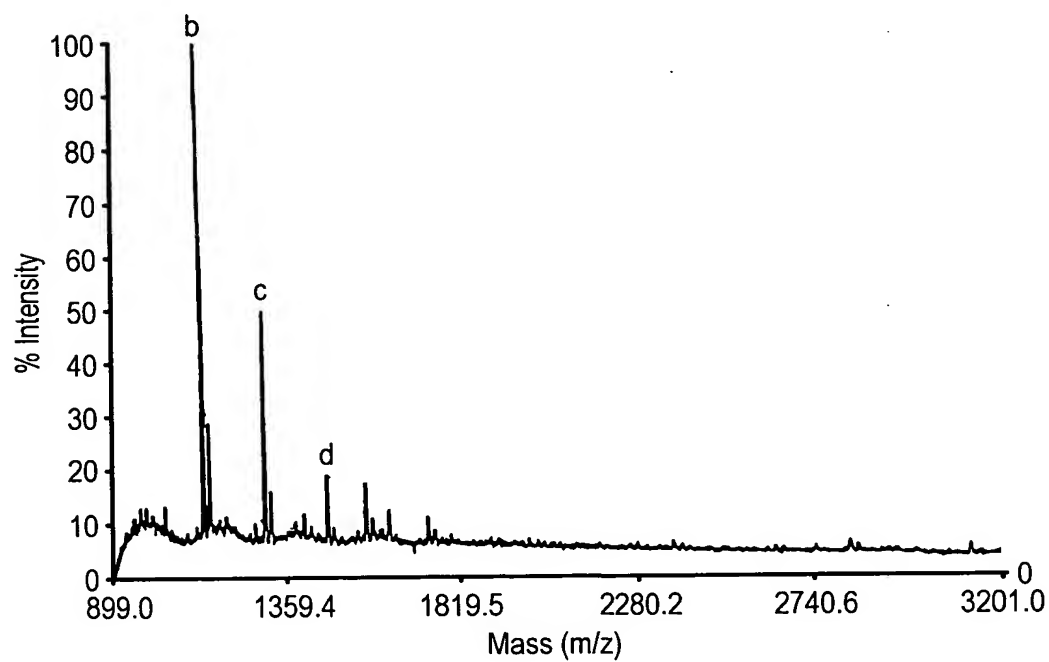
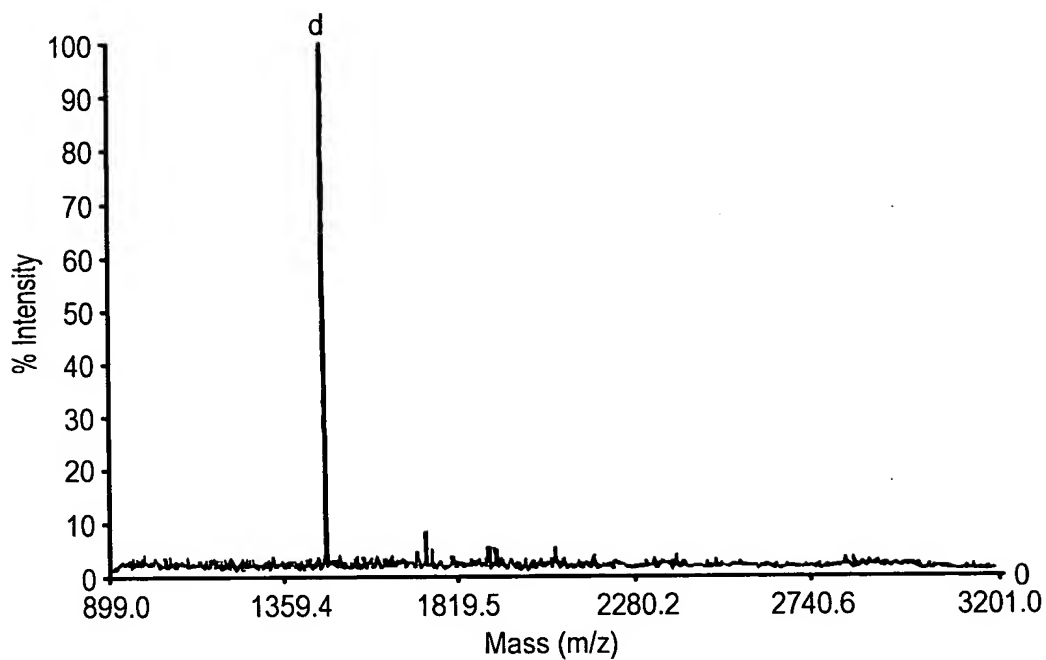
ATGAAGATGAGACGCTACAAGCTCTTTCTCATGTTCTGTATGGCTGGCCTGTGCCTCATATCCTTCCTGC  
ACTTCTTTAAGACCTTATCCTATGTACCTTCCCGAGAGAACTGGCCTCCCTCAGCCCTAACCTCGTATC  
CAGCTTCTTCTGGAACAATGCCCCTGTCACTCCCCAGGCCAGTCCGGAGCCGGGTGGCCCCGACCTATTG  
CGGACACCCCTCTACTCCCACTCTCCCCTGCTCCAGCCACTGTCCCCGAGCAAGGCCACAGAGGAACTGC  
ACCGGGTGGACTTCGTGTTGCCGGAGGACACCACGGAGTATTTTGTGCGCACCAAAGCTGGTGGTGTGTG  
CTTCAAACCAGGTACCAGGATGCTGGAGAAACCTTCGCCAGGGCGGACAGAGGAGAAGCCCGAAGTGTCT  
GAGGGCTCCTCAGCCCGGGGACCTGCTCGGAGGCCCATGAGGCACGTGTTGAGTACGCGGGAGCGCCTGG  
GCAGCCGGGGCACTAGGCGCAAGTGGGTTGAGTGTGTGTGCCTGCCAGGCTGGCACGGGCCCAGTTGCGG  
GGTGCCCACGGTGGTGCAGTATTCCAACCTGCCACCAAGGAACGCCTGGTACCAGGGAGGTACCGAGG  
CGGGTTATCAACGCCATCAACATCAACACAGAGTTTCGACCTGCTGGATGTGCGCTTCATGAGCTGGGAG  
ATGTTGTGGACGCCCTTCGTGGTCTGTGAATCTAATTTACCGCCTACGGGGAGCCTCGGCCGCTCAAGTT  
CCGAGAGATGCTGACCAATGGCACCTTCGAGTACATCCGCCACAAGGTGCTCTATGTCTTCTCGGACCAT  
TTCCACCTGGTGGCCGTGAGGACGGCTGGATTGCGGATGACTACCTGCGCACCTTCCTCAGCCAGGATG  
GCGTCTCCCGCCTGCGCAACCTGCGGGCCGATGACGTCTTTATCATCGACGATGCGGACGAGATCCCTGC  
GCGTGATGGTGTGCTGTTCTCTCAAACTCTACGATGGCTGGACAGAGCCCTTCGCCTTCACATGCGGAAG  
TCCCTGTATGGTTTCTTCTGGAAGCAGCCGGGCACTGGAGGTGGTGTGAGGCTGCACCATGGACATGC  
TGCAGGCCGTGTATGGGCTGGATGGCATCCGCCTGCGCCGCGCCAGTACTACACCATGCCCAACTCCG  
GCAGTATGAGAACCGCACCGGCCACATCCTAGTGACGTGGTCTCTCGGCAGCCCCCTGCACCTTCGCGGGC  
TGGCATTGCTCCTGGTGTCTCACACCCGAGGGCATCTACTTTAAACTCGTGTGAGCCCAAGATGGCGACT  
TCCCCCGCTGGGGTGACTATGAGGACAAGAGGGACCTCAATTACATCCGCAGCTTGATCCGCAGTGGGGG  
ATGGTTTCGACGGAACGCAGCAGGAGTACCCTCCTGCGGACCCAGTGAGCACATGTATGCTCCTAAATAC  
CTGCTCAAGAACTATGACCAGTTCGCTACTTGTGGAATAATCCCTACCGGGAGCCCAAGAGCACTGTAG  
AGGGTGGGCGCCAGAACCAGGGCTCAGATGGAAGGCCATCTGCTGTGAGGGGCAAGTTGGATACAGTGA  
GGGCTAG

>gi|2117717|pir||JC4362 beta-1,4-mannosyl-glycoprotein 4-beta-N-  
acetylglucosaminyltransferase (EC 2.4.1.144) III - mouse  
MRRYKLFMFCMAGLCLISFLHFFKTLSTVTFPRELASLSPNLISFFWNNAPVTPQASPEPGDPDLLRT  
PLYSHSPLLQPLSPSKATEELHRVDFVLPEDTTEYFVRTKAGGVCFKPGTRMLEKPSPGRTEEKTEVSEG  
SSARGPARRPMRHLVSSRERLGSRGTRRKWVECVCLPGWHGPPSCGVPTVVQYSNLPTKERLVPREVPRRV  
INAININHEFDLLDVRFHLDVDFVCDNFTAYGEPRLKPREMLTNGTFEYIRHKVLYVFLDHFP  
PGGRQDGIADDYLRFTLTQDGVSRRLNLRPDDVFIIDDADEIPARDGVLFLKLYDGTWTEPFAFHMRSLSL  
YGFfwKQPGTLEVVSCTMDMLQAVYGLDGIRLRRRQYYTTPNFQYENRTGHILVQWSLGSPLHFAGWH  
CSWCFTPEGIYFKLVSAQNGDFPRWGDYEDKRDNLNIRSLIRTGGWFDGTQOEYPPADPSEHMYAPKYL  
KNYDQFRYLLNPPYREPSTVEGGRQNQGSDDRSSAVRGKLDTAEG

FIG. 24

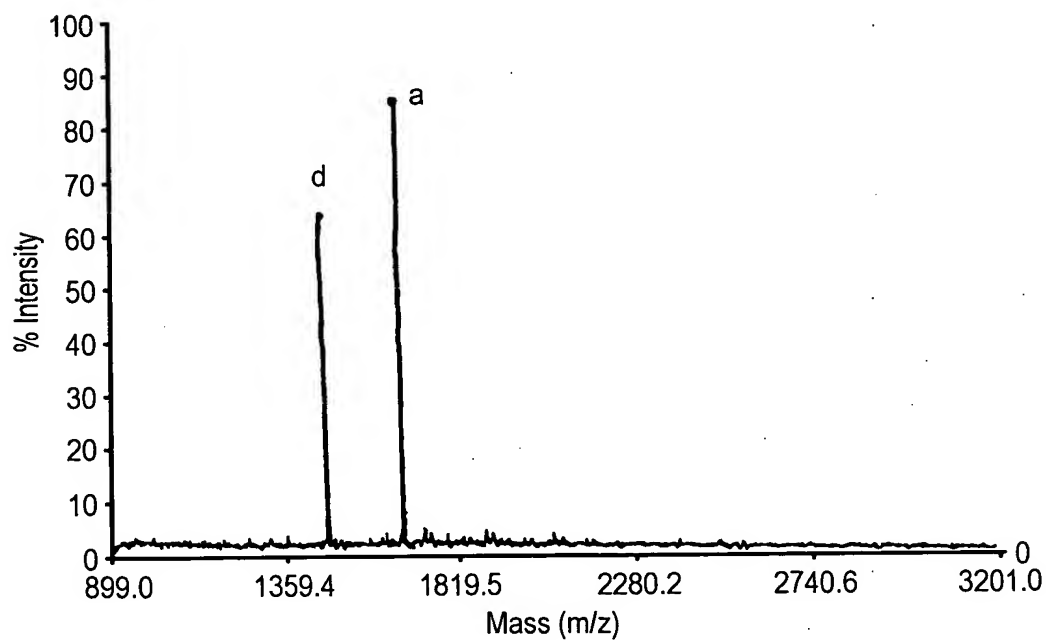
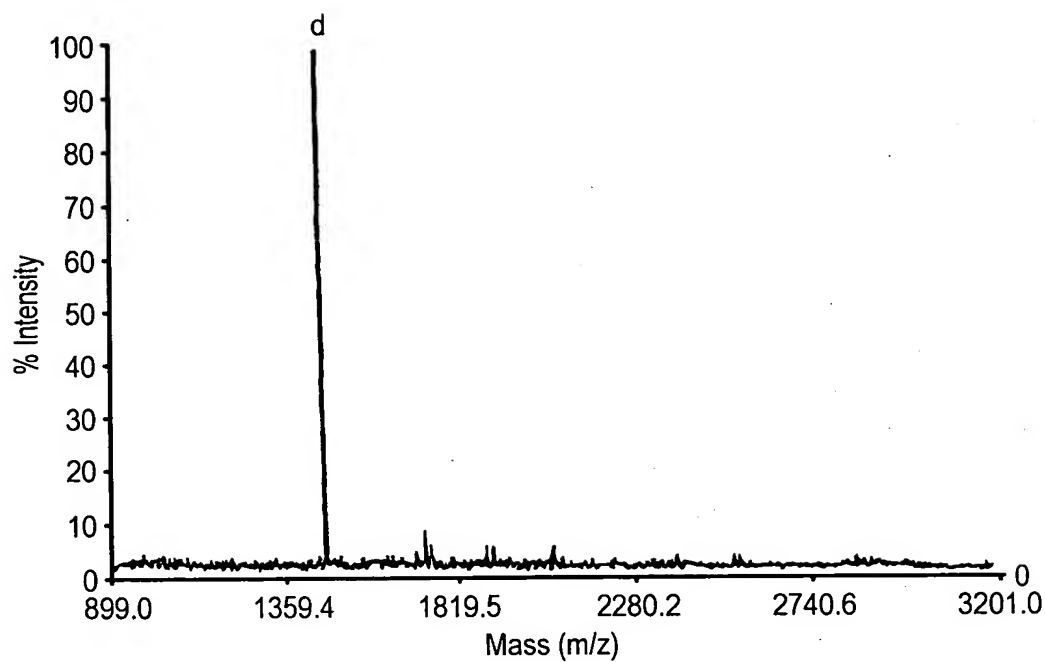
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FIG. 25



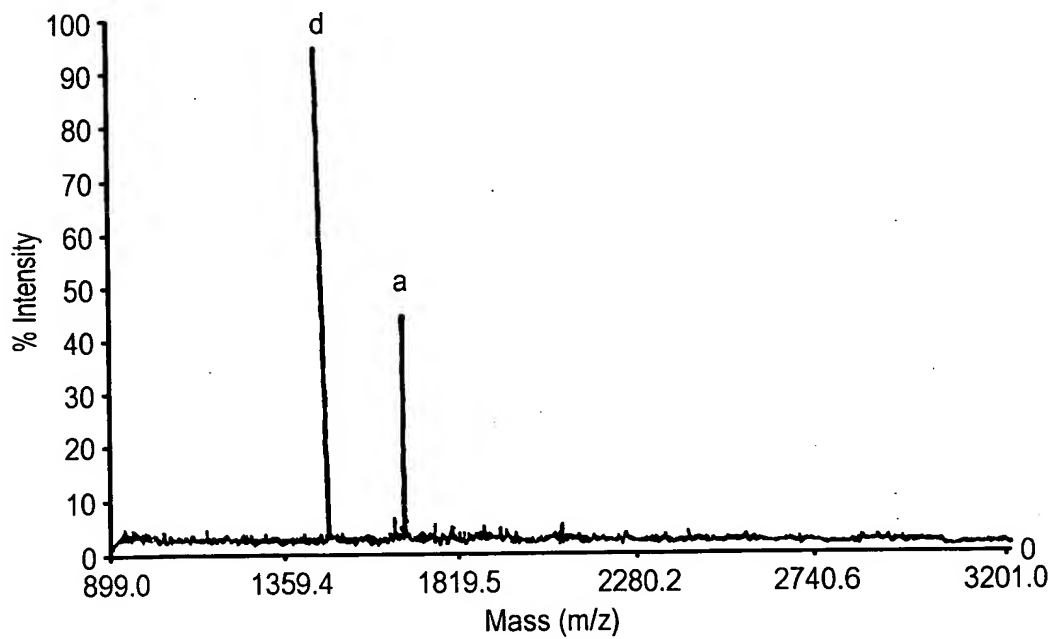
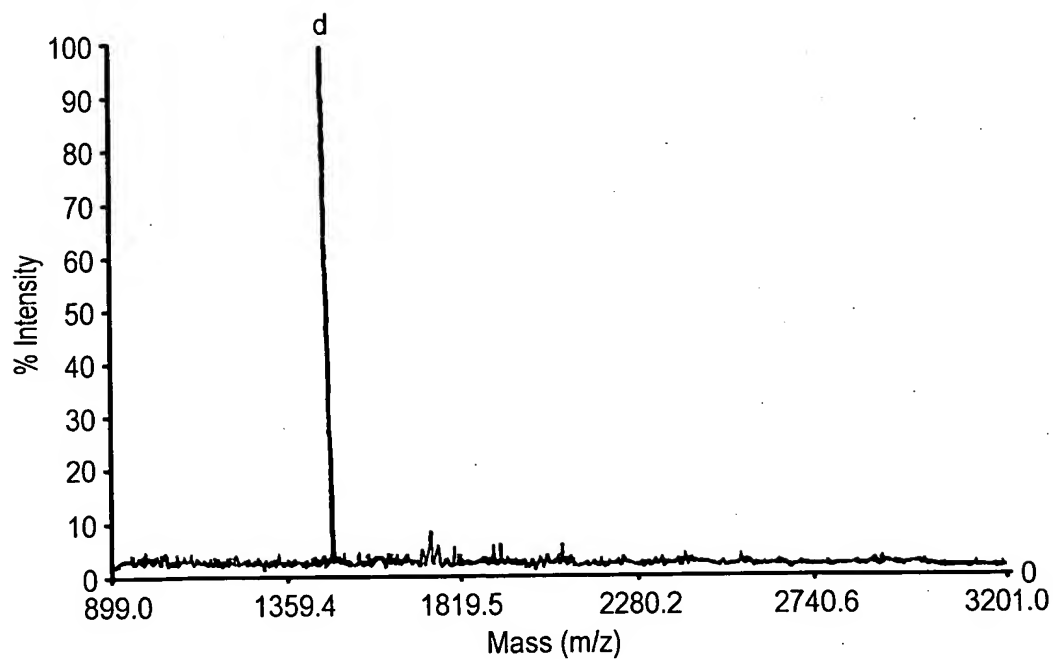
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FIG. 26



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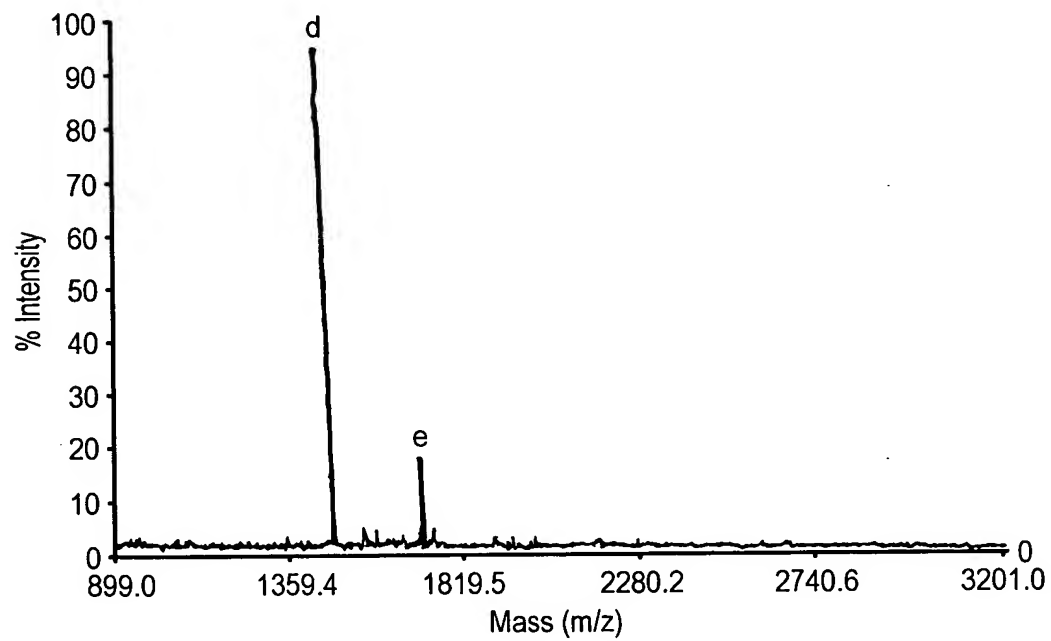
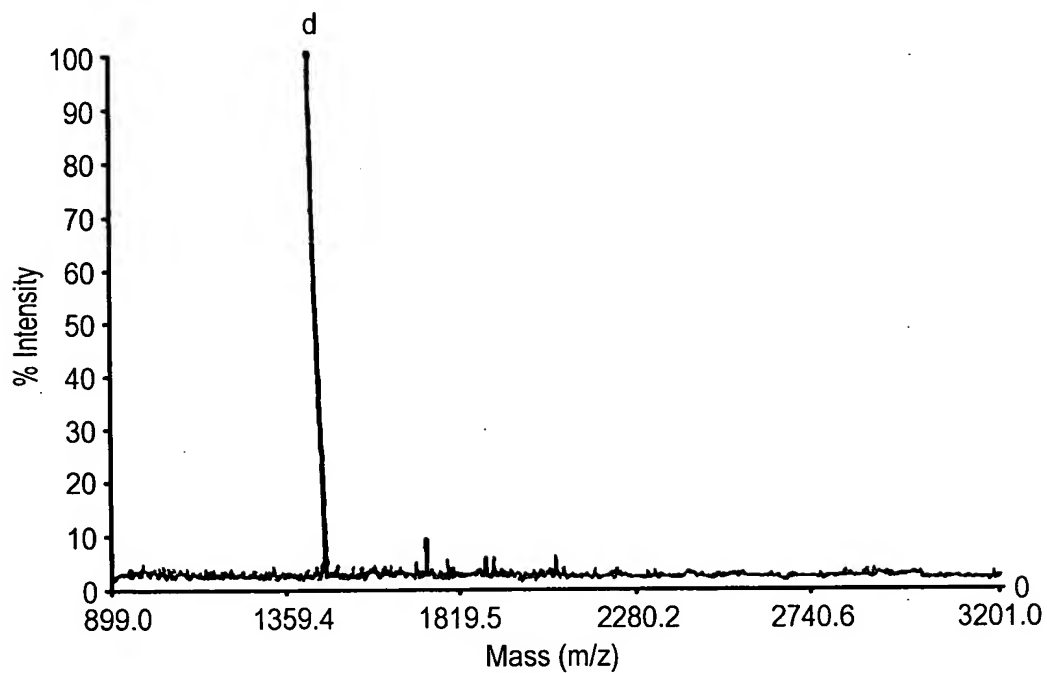
FIG. 27





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FIG. 28



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FIG. 29

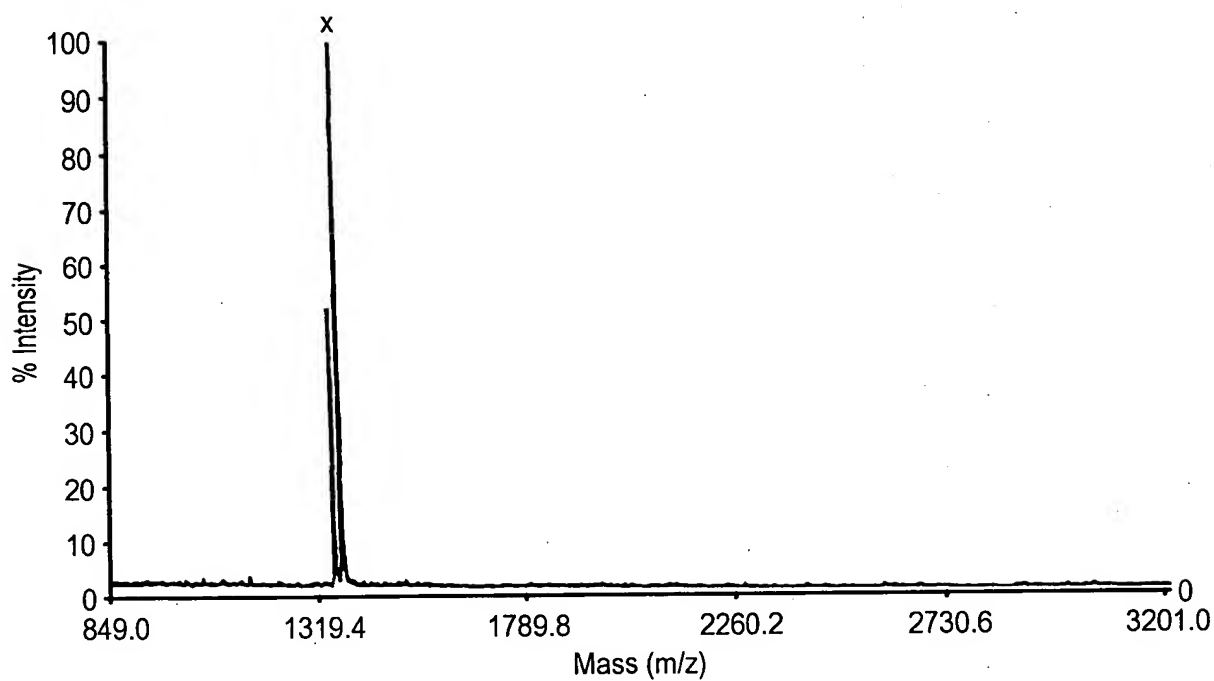
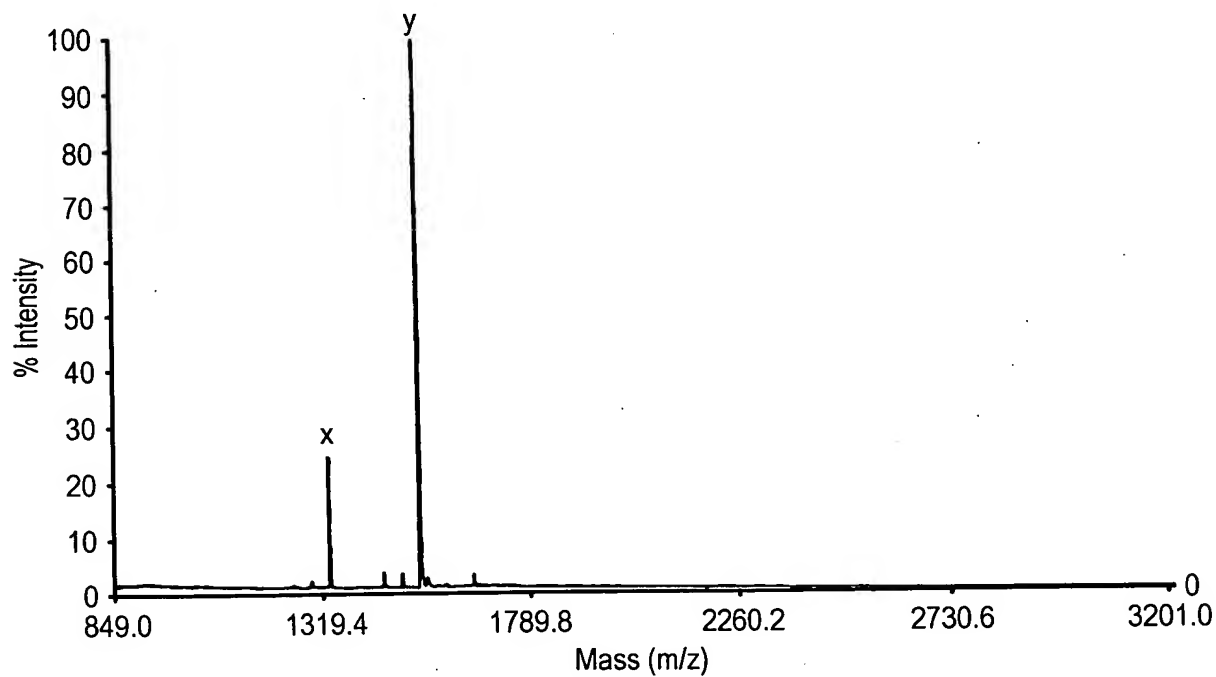


FIG. 30



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FIG. 31

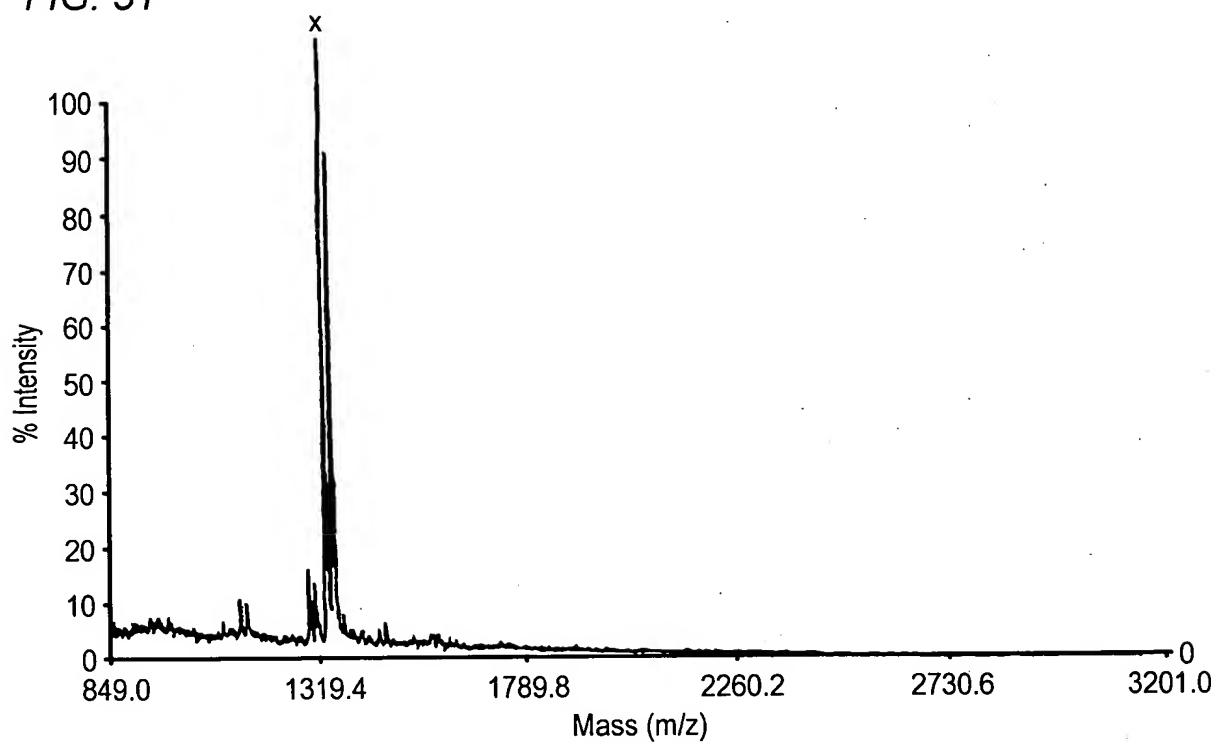
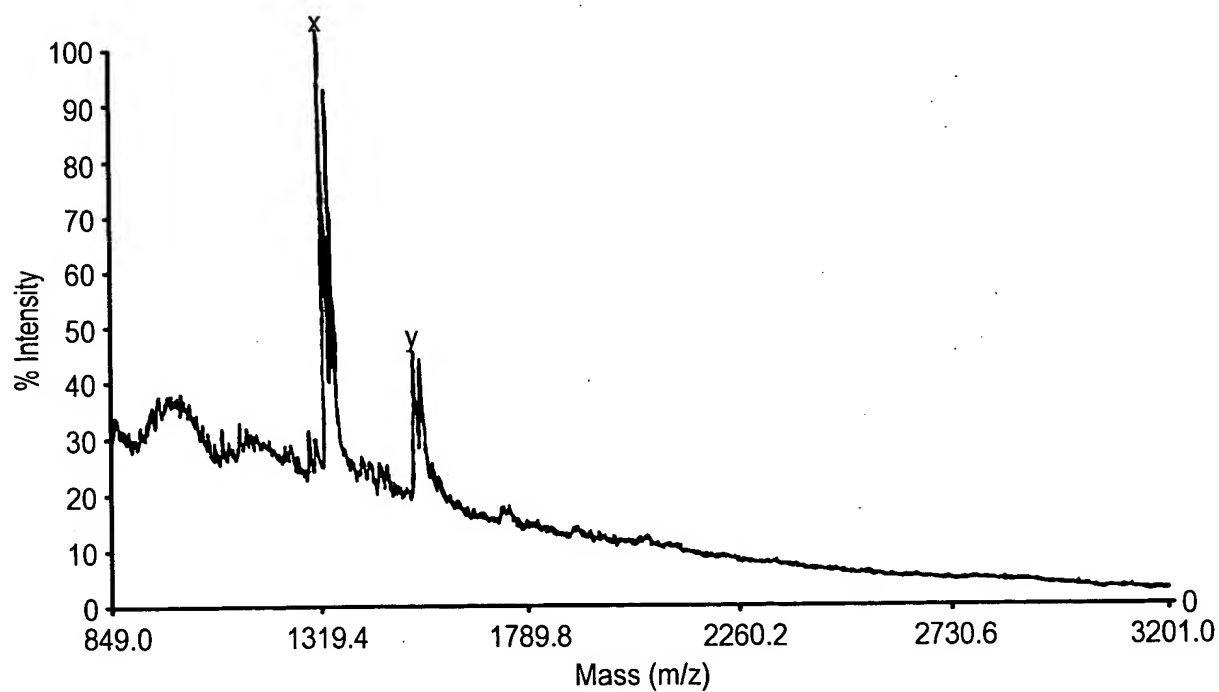
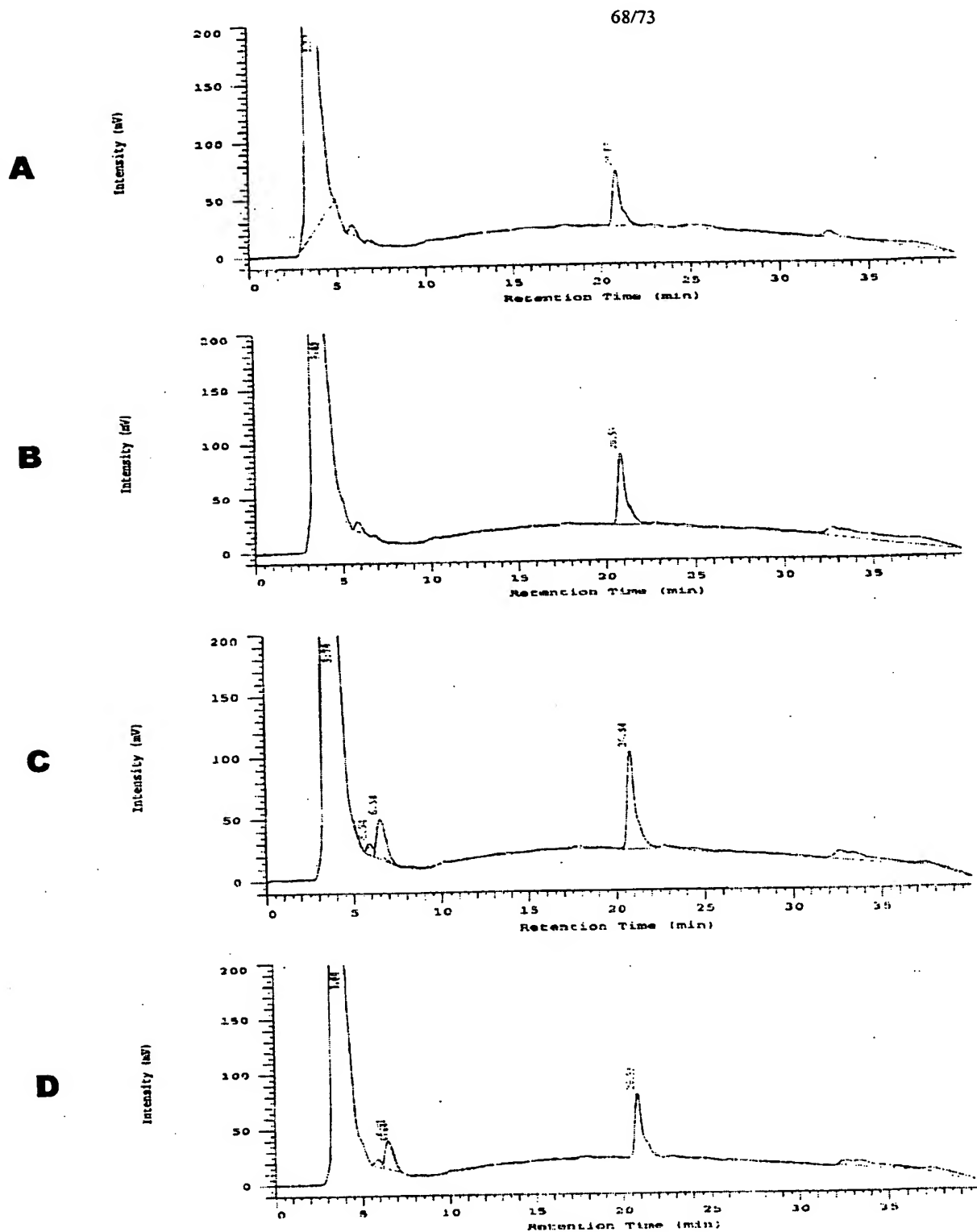


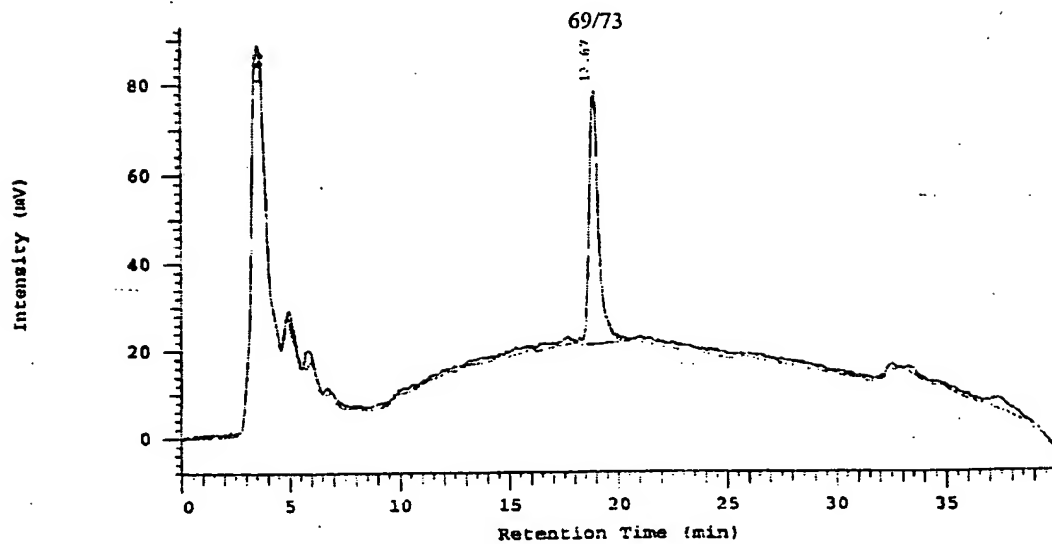
FIG. 32



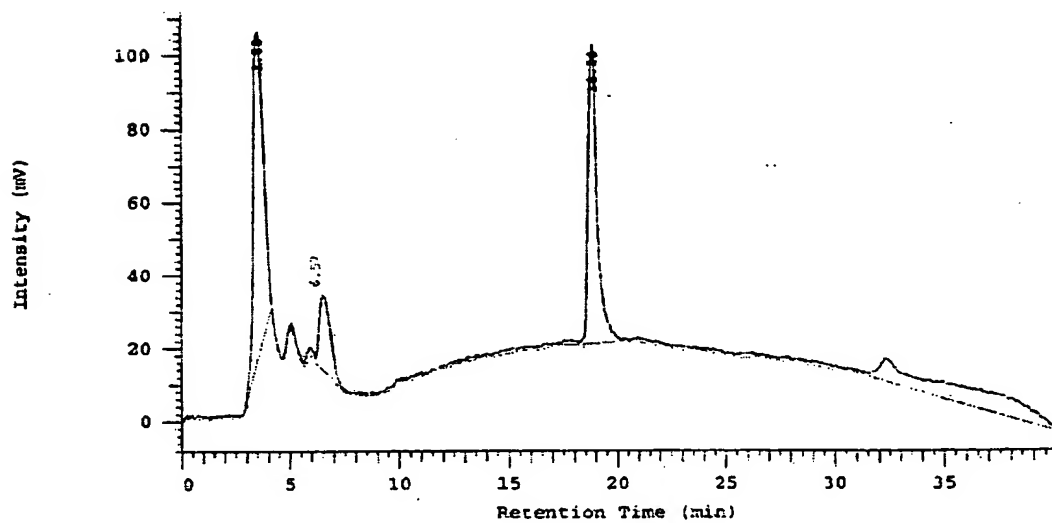


**Fig. 33**

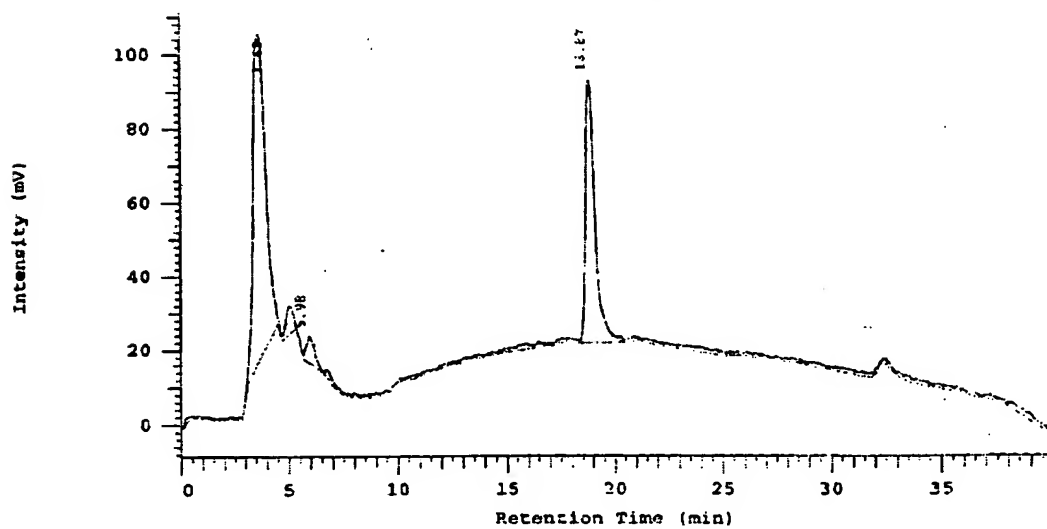
**A**



**B**

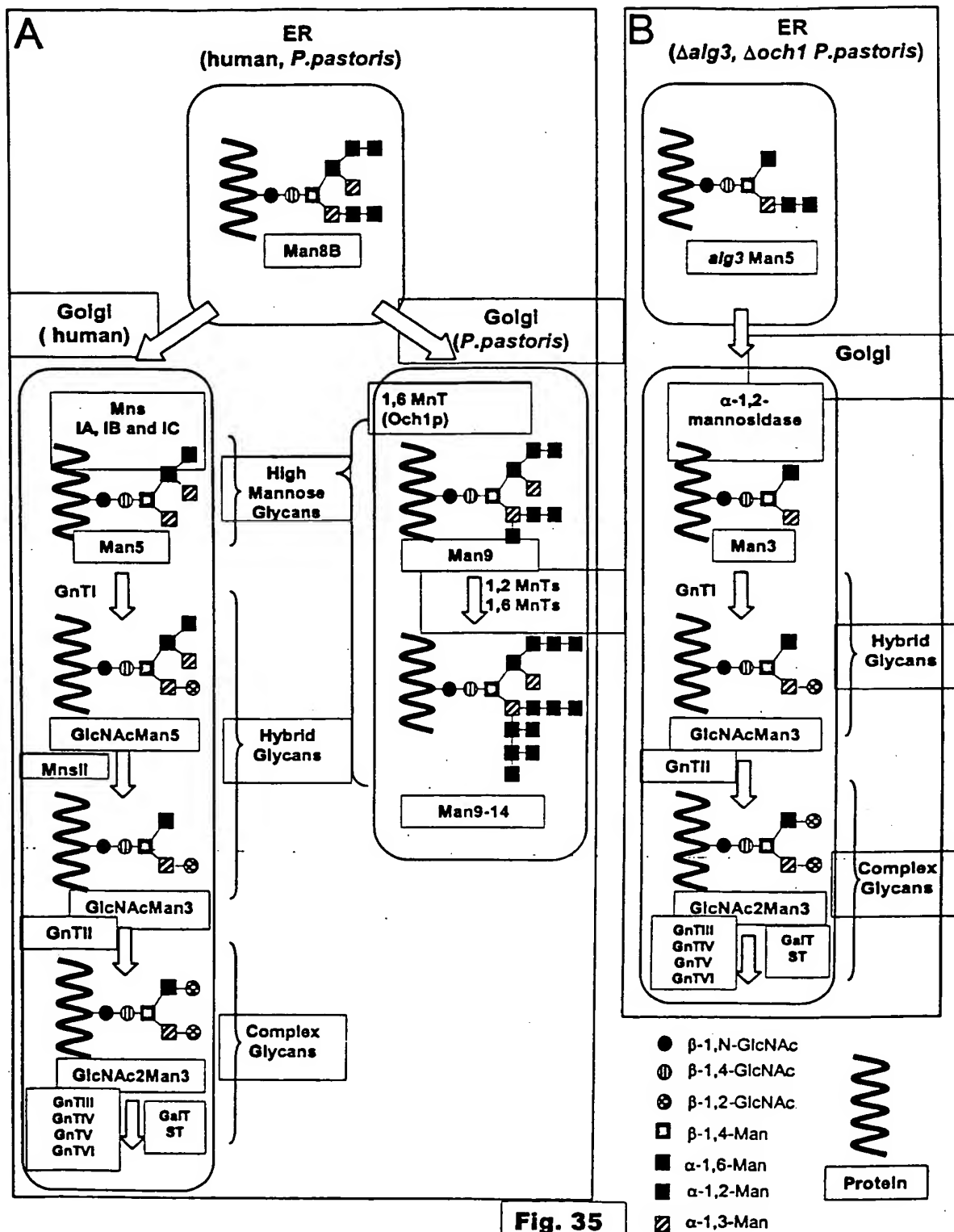


**C**



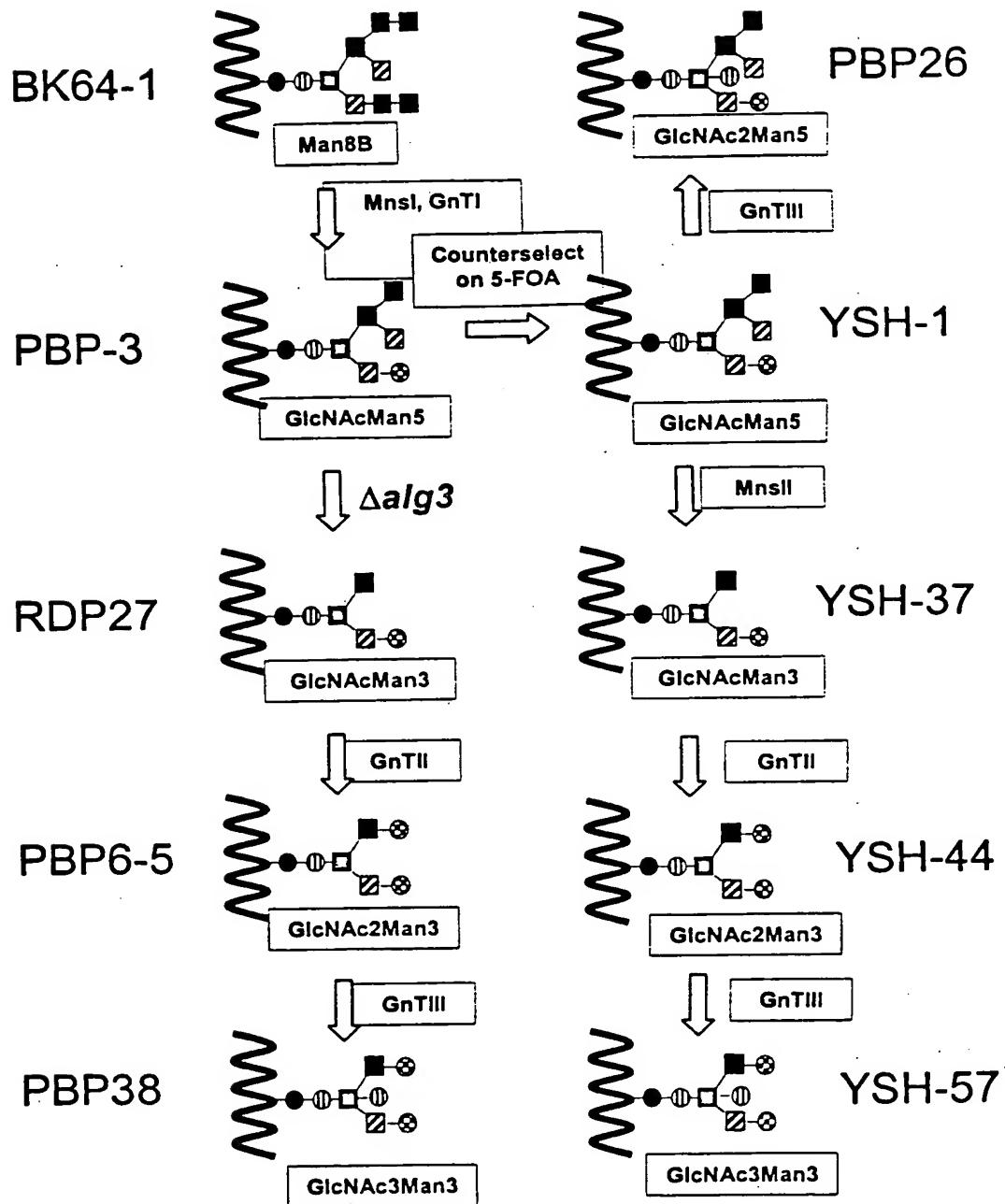
**Fig. 34**

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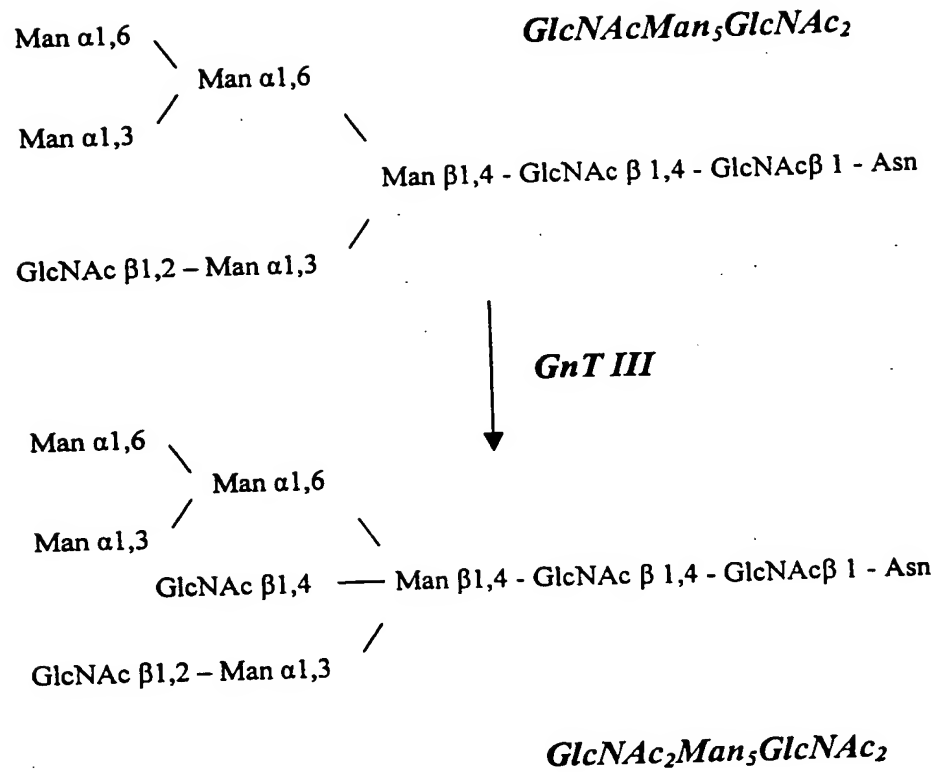
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**Fig. 36**



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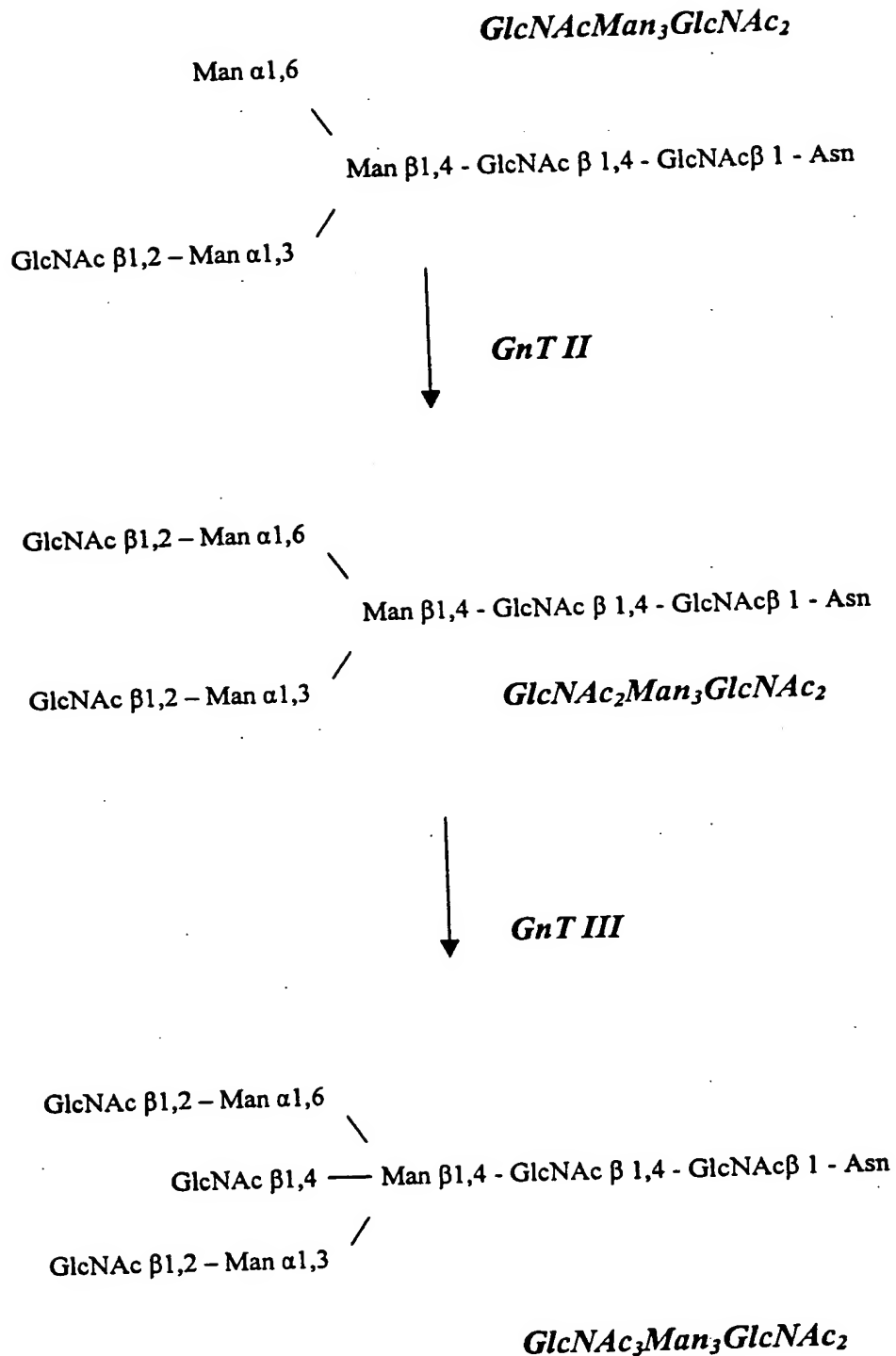
**Fig. 37**





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**Fig. 38**



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